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(54) Title: DETECTION OF MOLECULAR INTERACTIONS BY REPORTER SUBUNIT COMPLEMENTATION

(57) Abstract

Methods and compositions for detecting molecular interactions, particularly protein-protein interactions, are provided. The invention allows detection of such interactions in living cells or *in vitro*. Detection of molecular interactions in living cells is not limited to the nuclear compartment, but can be accomplished in the cytoplasm, cell surface, organelles, or between these entities. In one embodiment, the method utilizes novel compositions comprising fusion proteins between the molecules of interest and two or more inactive, weakly-complementing β -galactosidase mutants. Association between the molecules of interest brings the complementing β -galactosidase mutants into proximity so that complementation occurs and active β -galactosidase is produced. The active β -galactosidase may be detected by methods well-known in the art. Among the uses of the invention are the study of protein-protein interactions, functional genomics, agonist and antagonist screening and drug discovery.

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WO 98/44350 PCT/US98/06648

DETECTION OF MOLECULAR INTERACTIONS BY REPORTER SUBUNIT COMPLEMENTATION

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority to U.S. Provisional Patent Applications Serial No. 60/042,576 filed April 2, 1997 and Serial No. 60/054,638, filed August 4, 1997; and U.S. Patent Application Attorney Docket No. 28600-20206.00, filed April 1, 1998.

STATEMENT OF RIGHTS TO INVENTIONS MADE UNDER FEDERALLY SPONSORED RESEARCH

Not applicable.

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TECHNICAL FIELD

This invention is in the field of molecular biology and, more specifically, in the field of reporter systems useful for the analysis of protein-protein interactions.

BACKGROUND

The β-galactosidase enzyme (β-gal), the protein product of the *E. coli lacZ* gene, is widely used in studies of gene expression and cell lineage in higher organisms. Several biochemical assays of β-gal activity, including live-cell flow cytometry and histochemical staining with the chromogenic substrate 5-bromo-4-chloro-3-indolyl β-D-galactopyranoside (X-gal) make the product of the *lacZ* gene extremely versatile as a quantitative reporter enzyme, selectable marker, or histological indicator. Bronstein *et al.* (1989) *J. Biolumin. Chemilumin.* 4:99-111; Nolan *et al.* (1988) *Proc. Natl. Acad. Sci. USA* 85:2603-2607; and Lojda (1979) *Enzyme Histochemistry: A Laboratory Manual*, Springer, Berlin. One property of the *lacZ* system that has been well-characterized in studies of bacterial genetics, but has not been exploited in eukaryotes is the phenomenon of intracistronic complementation. Studies in *E. coli* have shown that deletions of β-gal which remove portions of either the N-terminus or the C-terminus produce enzyme which is inactive. However, coexpression of one of these deletion mutants with a second inactive deletion mutant containing domains that are lacking in the first can restore β-gal enzymatic activity in a process called complementation. This complemented β-gal activity arises by

concentration-dependent assembly of a stable hetero-octameric enzyme complex comprising all the essential domains of the wild-type homotetramer. Ullman et al. (1965) J. Mol. Biol. 12:918-923; Ullman et al. (1967) J. Mol. Biol. 24:339-343; and Ullman et al. (1968) J. Mol. Biol. 32:1-13.

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A system utilizing β -gal complementation in enzyme assays has been described. Henderson, U.S. Patent 4,708,929. In this system, enzymatically inactive β -gal polypeptide fragments, capable of combining with high affinity to form active β -gal by complementation, are used. One of the fragments is conjugated to analyte, which allows it to compete with analyte for binding to an analyte-binding protein. If bound to the analyte-binding protein, the β -gal fragment is unable to complement. Thus, by comparing β -gal activity in the presence of sample to that obtained in the presence of a known concentration of analyte (at equal concentrations of analyte-binding protein) the amount of analyte in the sample may be determined. This method requires high-affinity complementing subunits of β -gal, requires that an analyte-binding protein be known, and is not applicable to single-cell analysis.

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Previous systems for the study of protein-protein interactions have been described which utilize two fusion genes whose products reconstitute the function of a transcriptional activator. Fields et al., (1989) Nature 340:245-247; Bai et al., (1996) Meth. Enzymol. 273:331-347; Luo et al., (1997) BioTechniques 22(2):350-352. In one fusion gene, a sequence encoding a first protein is conjugated to a sequence encoding a DNA-binding domain of a transcriptional regulatory protein. In a second fusion gene, a sequence encoding a second protein is conjugated to a sequence encoding a transcriptional activation domain of a transcriptional regulatory protein. The two fusion genes are co-transfected into a cell which also contains a reporter gene whose expression is controlled by a DNA regulatory sequence that is bound by the DNA-binding domain encoded by the first fusion gene. Expression of the reporter gene requires that a transcriptional activation domain be brought adjacent to the DNA regulatory sequence. Binding of the first protein to the second protein will bring the transcriptional activation domain encoded by the second fusion gene into proximity with the DNA-binding domain encoded by the first fusion gene, thereby stimulating transcription of the reporter gene. Thus, the level of expression of the reporter gene will reflect the degree of binding between the first and second proteins.

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There are several disadvantages associated with the use of the above-mentioned system. As it is dependent upon transcriptionally-regulated expression of a reporter gene, this system is limited to the assay of interactions that take place in the nucleus. In addition, the assay is indirect, relying on transcriptional activation of a reporter gene whose product is diffusible. Hence, a method which would allow a direct and immediate examination of molecular interactions, at the site where they occur, would be desirable.

A system for detecting protein-protein interactions, not limited to nuclear interactions, has been described. U.S. Patent Nos. 5,503,977 and 5,585,245. In this system, fusions between potential interacting polypeptides and mutant subunits of the protein ubiquitin are formed. Juxtaposition of the two ubiquitin subunits brought about by interaction between potential interacting polypeptides creates a substrate for a ubiquitin-specific protease, and a small peptide reporter fragment is released. In this system, binding between the potential interacting polypeptides does not generate any type of enzymatic activity; therefore, signal amplification is not possible. Additionally, the ubiquitin system does not measure activity in intact cells, but relies on assays of proteolysis in cell-free extracts. What is needed is a sensitive method for examining protein interactions in intact cells in the relevant cellular compartment.

Fluorescence imaging has been used to study the intracellular biochemistry of living cells. A fluorescent indicator for the adenosine 3',5'-cyclic monophosphate (cAMP) signaling pathway has been described in which the sensor is a cAMP kinase in which the catalytic and regulatory subunits each are labeled with a different fluorescent dye, such as fluorescein or rhodamine, capable of fluorescence resonance energy transfer in the holoenzyme complex. A change in shape of the fluorescence emission spectrum occurs upon cAMP binding, and therefore activation of the kinase can be visualized in cells microinjected with the labeled holoenzyme. Adams et al., Nature, 349: 694-697 (1991). This system is limited by the fact that it requires microinjection, and a preferred distance between the labeled units for energy transfer to occur.

Substrates for β -lactamase have been described in the art which include a fluorescent donor moiety and a quencher, which include an attached group which makes them permeable through cell membranes, wherein the attached group is hydrolyzed off after the substrate enters the cell. Fluorescence energy transfer between the donor and quencher is monitored as an indicator of β -lactamase activity. This system also can be

used in a reporter gene assay using cells containing β -lactamase reporter genes functionally linked to a promoter. PCT WO 96/30540 published October 3, 1996, the disclosure of which is incorporated herein.

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DISCLOSURE OF THE INVENTION

The present invention provides methods and compositions for detecting, assaying and quantitating molecular interactions within living cells and *in vitro*, through complementation between two or more low affinity reporter subunits, such as distinct *E. coli lacZ* mutations. In a preferred embodiment, protein-protein interactions within living cells are detected and quantitated using the methods and compositions of the present invention. The practice of the present invention enables, for the first time, the study of protein-protein interactions and their control in living mammalian cells without reliance upon the transcriptional activation of a reporter gene construct. Association of the proteins of interest results directly in enzyme activity and is independent of other cellular functions. Therefore, the present invention provides advantages over other systems currently in use by allowing the detection of complexes that are excluded from the nucleus, and detection of complexes whose formation would inhibit transcription. Furthermore, the present invention allows the detection and localization of specific binding interactions within cells at different stages of development and differentiation, and an analysis of the induction or inhibition of binding interactions in cells.

Interactions occurring within the nucleus of the cell, interactions occurring in the cytoplasm, on the cell surface, within or on the surface of organelles, or between cytoplasmic and surface (either cellular or organellar) molecules, as well a interactions occurring outside the cell, are all capable of being detected in the practice of the present invention. Thus, the invention surmounts the limitations associated with previous assays for protein-protein interactions, which were either limited to interactions occurring in the nucleus, or did not always allow accurate localization of molecular interactions, and which were not well-suited for detection of interactions which resulted in inhibition of transcription or translation.

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Accordingly, in one embodiment, the invention provides a reporter system component comprising:

a first low-affinity reporter subunit, coupled to a first putative binding moiety;

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wherein the first low-affinity reporter subunit is capable of association with at least a second low-affinity reporter subunit to generate a detectable signal, said association being mediated by the first putative binding moiety.

In another embodiment, the invention provides a method of determining the occurrence of binding between first and second putative binding moieties, the method comprising:

providing a reporter system comprising: a)

a first component comprising a first low affinity reporter subunit, coupled to the first putative binding moiety; and

a second component comprising a second low affinity reporter subunit coupled to the second putative binding moiety;

wherein the first low affinity reporter subunit is capable of association with and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal, said and the second low affinity reporter subunit to generate a detectable signal and the second low affinity reporter subunit to generate a detectable signal and the second low affinity reporter subunit to generate a detectable signal and the second low affinity reporter subunit to generate a detectable signal and the second low affinity reporter subunit to generate a detectable signal and the second low affinition and the association being mediated by the binding of the first and second putative binding moieties;

- combining the first component and the second component; and b)
 - detecting the presence or absence of the signal. c)

In a further embodiment, the invention provides a method of screening for binding of a first binding moiety with members of a plurality of different second putative binding moieties, the method comprising:

> providing a plurality of reporter systems each comprising: a)

a first component comprising a first low affinity reporter subunit coupled to the first binding moiety, and

one of a plurality of second components each comprising a second low affinity reporter subunit coupled to one of said plurality of second putative binding moieties, wherein in each of said second components, said second putative binding moiety is different;

wherein the first low affinity reporter subunit is capable of association with the second low affinity reporter subunit to generate a detectable signal upon the binding of the first binding moiety with one of said different second putative binding moieties;

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- b) individually combining the first component with each of the plurality of second components to produce a plurality of binding assay samples, each of which includes the first component and a different one of the second components; and
- detecting the presence or absence of the signal in each of the binding assay c) samples.

The invention additionally provides nucleic acids encoding fusion proteins including a low affinity reporter subunit and a putative binding moiety, and the fusion proteins encoded by said nucleic acids. The invention further provides viral vectors comprising nucleic acids encoding such fusions proteins. The invention also provides cells transformed by the nucleic acids and viral vectors described above.

All patents, patent applications and publications referred to herein are hereby incorporated by reference.

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Figure 1 is a schematic illustration of three deletion mutant lacZ constructs, 15 designated $\Delta\alpha$, $\Delta\omega$ and $\Delta\mu$.

> Figure 2A is a schematic illustration of a viral construct encoding fusion proteins of the $\Delta\alpha$ or $\Delta\omega$ β -gal mutants with either the intracellular FKBP-rapamycin associated $\frac{1}{2}$ $\frac{$ protein (FRAP) or the intracellular rapamycin binding protein, FK506-binding protein-12 (FKBP12) upstream of the hygromycin or neomycin resistance genes.

Figure 2B is a schematic illustration of a viral construct encoding fusion proteins of the $\Delta\alpha$ or $\Delta\omega$ β -gal mutants with either FRAP or FKBP12 and another protein, represented as x and x', upstream of the hygromycin or neomycin resistance genes.

Figures 3A and 3B show X-gal staining of fixed cells expressing both FKBP12- $\Delta\omega$ and FRAP- $\Delta\alpha$. Cells shown in 3b were exposed to 10 ng/ml rapamycin for 12 hr. Cells shown in 3a were not exposed to rapamycin.

Figure 4A is a graph of β -gal activity vs. time with and without rapamycin treatment of C2C12 cells expressing both FKBP12- $\Delta\omega$ and FRAP- $\Delta\alpha$ fusion proteins.

Figure 4B is a graph of the dose-response to rapamycin of the activity of β -gal in C2C12 cells expressing both FKBP12-Δω and FRAP-Δα fusion proteins.

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Figure 5 shows rapamycin-dependent increase in β -gal activity in lysates from cells expressing both FKBP12- $\Delta\omega$ and FRAP- $\Delta\alpha$ fusion proteins, measured by chemiluminescence.

Figure 6A shows analysis by Fluorescence-Activated Cell Sorting (FACS) of C2C12 cells expressing both FKBP12- $\Delta\omega$ and FRAP- $\Delta\alpha$ after 90 minutes of rapamycin treatment. Dark peaks represent profiles obtained from untreated samples; light peaks represent profiles from samples that have been treated with 10 ng/ml rapamycin.

Figure 6B shows a FACS profile of untreated cells and indicates a subpopulation selected on the basis of low β -gal activity.

Figure 6C shows FACS analysis of the subpopulation of cells selected in Figure 6B after overnight culture in the absence (dark peak) or presence (light peak) of rapamycin.

In Figures 6A, 6B and 6C, the vertical axis represents cell number and the horizontal axis represents intensity of β-gal fluorescence expressed on a logarithmic scale.

Figure 7 shows EGF receptor dimerization monitored using β -gal $(\beta, \beta, \beta, \beta) = (\beta, \beta, \beta) + (\beta, \beta, \beta$

Figure 7A depicts schematically the rationale of the assay: two weakly complementing deletion mutants of β -gal are linked to the extracellular and transmembrane domains of the EGF receptor. Receptor dimerization, stabilized by EGF; will drive β -gal complementation.

Figure 7B shows the design of the retroviral constructs used in the assay. *E. coli* lacZ deletion mutants $\Delta\alpha$ and $\Delta\omega$ were cloned into pWZL vectors expressing neomycin or hygromycin resistance, respectively. The extracellular and transmembrane (tm) domains of human EGF receptor were cloned in frame with the $\Delta\alpha$ and $\Delta\omega$ mutants.

Figure 7C shows FACS analysis of a population of transduced and selected cells. EGF treatment increases the β -gal activity (fluorescein fluorescence) in a substantial proportion of the cells. The FACS profile of cells without EGF treatment is shaded in light gray and is outlined in white. The profile of cells treated with EGF is shaded dark gray.

Figure 7D shows FACS analysis of chimeric receptor expression, using a monoclonal antibody to the extracellular domain of the human EGF receptor. The FACS profile of the transduced and selected population is shaded medium gray and outlined in white; untransduced cells are shaded light gray and outlined in white. The FACS was used to clone cells that had low β-gal activity in the absence of EGF and showed increased β-gal

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activity in the presence of EGF. One clone that had low levels of the chimeric receptor relative to the population (shaded in dark gray) was used for further analyses.

Figure 7E shows induction of EGF receptor dimerization (β -gal activity) in all of the cells of the clone selected in Figure 7D, upon treatment with 100 ng/ml EGF for two hours. Untreated cells are shaded in light gray and outlined in white; EGF treated cells are shaded in dark gray.

Figure 7F shows that dimerization can be detected after very short treatments with EGF. Cells were treated with 100 ng/ml EGF for 0, 1, 4, 8, and 15 minutes before cells were rinsed and processed for FACS analysis. The mean fluorescence of the cell sample is plotted.

Figure 9 shows that EGF receptor dimerization is enhanced by tyrphostin AG1478.

Figure 9A shows, in the left panel, schematic diagrams of different regimens for treatment of cells with EGF, tyrphostin, or both. After the various treatments, cells were analyzed on the FACS, and the mean fluorescence is shown in the right panel. Each treatment was performed in triplicate.

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MODES FOR CARRYING OUT THE INVENTION

Definitions

As used herein, the following terms have the following definitions:

As used herein, a "reporter subunit" refers to a member of a complex of two or more subunits which are capable of associating with low binding affinity with each other to generate a detectable signal, or which are capable of associating with each other and one or more additional substances to generate a detectable signal, and which do not individually generate the detectable signal.

As used herein, "low affinity" reporter subunits refer to molecular species which have a sufficiently low binding affinity for each other such that when they each are covalently attached to two different binding moieties, they substantially do not become associated unless a binding interaction between the two binding moieties occurs. "Low affinity" thus generally refers to a binding affinity which is at least less than that of the 1900 to 100 t a part that the satisfied binding moieties. The apparatus of this object the mark and the open in the satisfied by

As used herein, "binding moieties" refers to at least two molecular species, such as proteins or fragments thereof, which interact with each other to form a stable complex.

As used herein, a "detectable signal" refers to any detectable signal which occurs which is a second the association of the reporter subunits or via the interaction of the associated and the second and the sec subunits with another substance. The detectable signal may be for example, a chromogenic, fluorescent, phosphorescent or chemiluminescent signal, such as a detectable product of an enzymatic reaction catalyzed by the associated reporter subunits.

The terms "protein", "polypeptide", and "peptide" are used interchangeably herein to refer to polymers of amino acids of any length. The polymer may be linear or branched, it may comprise modified amino acids, and it may be interrupted by non-amino acids. It also may be modified naturally or by intervention; for example, disulfide bond formation, glycosylation, myristylation, acetylation, alkylation, phosphorylation or dephosphorylation. Also included within the definition are polypeptides containing one or more analogs of an amino acid (including, for example, unnatural amino acids) as well as other modifications known in the art.

Unless otherwise indicated, the practice of the present invention will employ conventional techniques of molecular biology, biochemistry, microbiology, recombinant DNA, nucleic acid hybridization, genetics, immunology, embryology and oncology which

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are within the skill of the art. Such techniques are explained fully in the literature. See, for example, Maniatis, Fritsch & Sambrook, MOLECULAR CLONING: A LABORATORY MANUAL, Cold Spring Harbor Laboratory Press (1982); Sambrook, Fritsch & Maniatis. MOLECULAR CLONING: A LABORATORY MANUAL, Second Edition, Cold Spring Harbor Laboratory Press (1989); Ausubel, et al., CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons (1987, 1988, 1989, 1990, 1991, 1992. 1993, 1994, 1995, 1996).

Reporter Subunits

As used herein, a "reporter subunit" refers to a member of a complex of two or more subunits which are capable of associating with low binding affinity with each other to generate a detectable signal, or which are capable of associating with each other and one or more additional substances to generate a detectable signal, and which do not , এই মুক্ত কি individually generate the detectable signals কি এক প্রকরণে তেওঁকোনে মুক্ত কর্মকার এই ক্রিকালে কি সংগ্রহ

The detectable signal thus provides an indication that the subunits have become a second of the subunits have become associated. In general, in an assay of the binding affinity of a first and at least a second with the second secon and the second component is provided which are second component in the second component in the second component is provided which are second component in the second component in the second component is provided which are second component in the second component in the second component is provided which are second component in the second component in the second component component is provided which are second component component in the second component co includes one reporter subunit attached to the first molecular species, and a second with the second राह देखका component is provided which includes another of the same or different reporter subunit के अपने कार कर कर कर के attached to the second molecular species. The reporter subunits preferably have sufficiently low binding affinity for each other such that they substantially do not associate with each other in solution unless and until the molecules for which binding affinity is being assayed have sufficient binding affinity to mediate complex formation between the two components. Upon binding of the binding moieties and resulting association of the reporter subunits, generally by non-covalent interactions, such as hydrogen bonding or hydrophobic interactions, for example, the reporter subunits are oriented close enough to each other such that they are capable of associating with low affinity and generating a detectable signal. In the system, individual reporter subunits are not able to generate the detectable signal. Thus, the reporter subunits undergo forced complementation when brought into close proximity.

> The reporter subunits can be designed to have a preferred low affinity for a particular-application and for the conditions in which the binding assay is done. Binding of molecules will depend upon factors in solution such as pH, ionic strength, concentration

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of components of the assay, and temperature. In the binding assays using reporter systems described herein, the binding affinity of the reporter subunits should be low enough to permit forced complementation. Non-limiting examples of dissociation constants of the reporter subunits in an assay solution, such as a buffered system or cell interior, are on the order of greater than about 10⁻⁸ M for example, greater than 10⁻⁶ M or optionally, between about 10⁻² to 10⁻⁵ M depending upon the properties of the particular assay system.

Reporter subunits which have sufficiently low binding affinity, and yet are still capable of associating and generating a detectable signal upon the binding of molecular species attached to them can be designed as disclosed herein. Reporter subunits which can be used include any low binding affinity subunits which are capable of associating to produce a detectable signal. In one preferred embodiment, the reporter subunits are proteins which are capable of associating and are capable when associated of catalyzing a to an increase and increaction which produces a directly or indirectly detectable product. See the second of the line of the

na substrate to a detectable approach enzymes capable of catalyzing conversion of a substrate to a detectable approaches the conversion of a substrate to a substrate to a substrate the conversion of a substrate to a sub control 15 acres preaction product, either directly or indirectly, which have been used, for example, in cell and a care acres of the based screening assays may be used as reporter subunits. The enzymes can be modified as a session of the based ு நாகு காக்கையுள்ள complementation. These may be modified, for example, by site directed or random ாக வக்கையில் அவர mutagenesis, or deletion mutation, to provide low affinity subunits which are capable of associating with low binding affinity and thereby undergo complementation to catalyze an enzymatic reaction. For example, reporter subunits capable of complementation with low binding affinity may be derived from enzymes such as β -galactosidase, β -glucuronidase. (GUS), β-lactamase, alkaline phosphatase, peroxidase, chloramphenicol acetyltransferase (CAT) and luciferase. Any of a range of enzymes capable of producing a detectable product either directly or indirectly may be so modified or may occur naturally. Additionally, reporter subunits may be derived from non-enzymatic molecules. For example, association of two proteins may generate a unique conformation in one or both of the interacting proteins that can be recognized by an antibody or other ligand.

> β-galactosidase, which is encoded by the E. coli lacZ gene, is an enzyme which has been developed in the art as reporter enzyme. β-galactosidase activity may be measured by a range of methods including live-cell flow cytometry and histochemical staining with the chromogenic substrate 5-bromo-4-chloro-3-indolyl β-D-galactopyranoside (X-Gal). Nolan

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et al., Proc. Natl. Acad. Sci., USA, §5:2603-2607 (1988); and Lojda, Z., Enzyme Histochemistry: A Laboratory Manual, Springer, Berlin, (1979), the disclosures of which are incorporated herein.

Enzyme mutants capable of intracistronic complementation are especially suitable as reporter subunits. In E. coli, deletions of either the N or C terminus of β-gal produce enzyme that is inactive yet can be complemented by coexpression with a second inactive deletion mutant containing domains lacking in the first. The N- and C- terminal domains involved in complementation are known as the α and ω regions. Ullmann et al., J. Mol. Biol., 12:918-923 (1965); Ullman et al., J. Mol. Biol., 24:339-343 (1967); and Ullman et al. al., J. Mol. Biol., 32:1-13 (1968), the disclosures of which are incorporated herein. β-Gal. complementation systems in mammalian cells are described in Mohler and Blau, Proc. Natl. Acad. Sci. USA, 93:12423-12427 (1996), the disclosure of which is incorporated in the decision of the second holystelle, or the Macconstructed. A naturally occurring lacZ mutation, \(\Delta M\)! (Beckwith, \(J. Mol. Biol.) 8:427-47 (many constructed) 430 (1964); and Prentki, Gene, 122:231-232 (1992) and Nature, 369:761-766 (1994); the walkers in the same in the s 1940. The transmission of which are incorporated herein) designated as Δα herein may be constructed. The constructed the constructed of the const Another deletion mutation, designated Δω herein, was made as disclosed herein, and its structure is shown schematically in Figure 1. The peptide region between the or and w regions is referred to herein as the μ region, as first defined by Mohler and Blau, *Proc.* Natl. Acad. Sci. USA, 93:12423-12427 (1996). The $\Delta\alpha$ and $\Delta\omega$ mutants are demonstrated herein to have optimal forced complementation properties. These deletion mutants express polypeptides representing an α -acceptor/ ω -donor ($\Delta \alpha$) and an α -donor/ ω -acceptor ($\Delta \omega$).

> β-Gal complementation is based on the ability of mutant enzyme molecules to associate and reconstitute an active enzyme. Accordingly, two \u03b3-gal molecules that each lack one or more structural domains critical to the activity of the holoenzyme, associate to form a single functional unit that contains all of the required structural determinants. This phenomenon is dependent on the fact that interactions that would normally take place between domains of the single peptide of wild type β-gal, can also exist between domains present on two distinct peptides, leading to the formation of a stable dimer. This dimer behaves functionally as a single peptide of wild type β-gal, and participates ultimately in the formation of the tetramer that represents the active form of the enzyme. Thus, the ability of a pair of β-gal mutants to recreate an active form of the enzyme is strongly

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dependent on their ability to form a stable dimer and therefore would be expected to be dependent on their affinity for each other.

Surprisingly, it has been discovered that forced association or complementation of two distinct low affinity β-gal mutants results in an efficient formation of active enzyme molecules in mammalian cells even though they have relatively low affinity for each other. The forced complementation results when the two mutant subunits are brought into association due to the binding affinity of the binding moieties attached to the mutant subunits. By engineering constructs in which domains or proteins of interest drive the dimerization between $\Delta\alpha$ and

Δω β-gal mutants, it is possible to monitor and quantitate such interactions by assessing the efficiency of complementation obtained by coexpression of these fusion proteins in and the second intact eucaryotic cells.

ೂ ಎಂದು ರಾತ್ರಕ್ಷಣ ಹಿಳ್ಳಾಗಿ addition to two-component complementation between Δα and Δα β-gal mutants, ಆರು ಮಾಡುತ್ತಾರೆ ಅಂಗಾಗಿ ಆ The state of the invention also contemplates three-component complementation among mutants each of the state ் சார் வரு வருக்கிய முன்று a single functional a, யு or @ region. Among other applications; this வருக்கிய மாகி reporter. Similarly, higher-order systems containing four or more reporter components are with the components are epoch is a self-light within the scope of the invention. As the self-light s

> Using the fused protein systems, protein-protein interactions and their regulation can be studied in mammalian cells without relying on the transcriptional activation of a reporter construct. Association of the proteins of interest directly results in enzyme activity and is independent from other cellular functions. Therefore this system allows the detection of complexes that are excluded from the nucleus, or that involve partners that inhibit transcription. Furthermore it allows the detection, quantitation and determination of the localization of specific binding interactions within cells, as well as the temporal distribution of such binding interactions. Binding interactions may be compared in cells at different stages of development or differentiation, as well as in normal vs. pathologic cells and in infected vs. uninfected cells, to give but a few examples. Binding interactions can therefore be assessed against a background of endogenous competing components that may differ in nature and in concentration among different cell types.

Other enzymes may be identified or constructed which are capable of forced complementation in the reporter systems described herein. For example, the phenomenon

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of intracistronic complementation of enzymatic activity has been described for tryptophan synthetase. Jackson et al. J. Biol. Chem., 244:4539-4546 (1969). Complementation between mutant subunits of thymidylate synthase has been described. Pookanjanatavip et al., Biochemistry 31:10303-10309 (1992), the disclosure of which is incorporated herein. Thus, reporter subunits derived from any complementing enzyme system known in the art can be used in the practice of the present invention. Mutants can be derived from other enzymes or proteins that are capable of serving as reporters of protein-protein interactions, or whose activity can be regulated as described above. The system exploits the complementation ability of low binding affinity enzyme mutants for detection of protein-10 protein interactions.

For example, complementing low affinity reporter subunits derived from β-12 (2010). The first thing is lactamase can be constructed. Activity of the complementing β-lactamase can be detected சுராக நடிக்கு using substrates for β-lactamase developed in the art which include a fluorescent donor மாக மாக மக்கிய கடி the most see the moiety and a quencher, which include an attached group which makes them permeable and a quencher, which include an attached group which makes them permeable an attached group which makes the permeable at the permeable and the permeable at the permeable and the permeable at th 2年 1985年 1985 enters the cell. Fluorescence energy transfer between the donor and quencher; then can be 海绵 计模型 经营业 2010 1000 (1000) Semonitored as an indicator of β-lactamase activity, as described in PCT WO 96/30540 (1000) 基础 数据 数据 2000

In addition to enzymes which catalyze a reaction to produce a detectable product, proteins, protein domains or protein fragments which are themselves detectable upon association can be used. Exemplary proteins include green fluorescent proteins, which have characteristic detectable emission spectra, and have been modified to alter their emission spectra, as described in PCT WO 96/23810, the disclosure of which is incorporated herein. Fusions of green fluorescent proteins with other proteins, and DNA sequences encoding the fusion proteins which are expressed in cells are described in PCT WO 95/07463, the disclosure of which is incorporated herein.

Other exemplary subunits include subunits which are capable of associating to produce a photochemical signal such as a fluorescent or luminescent signal, including chemiluminescent or photoluminescent signals. The reporter subunits also may comprise fluorophores which are capable of detectable resonance energy transfer when they are closely associated, as disclosed, for example, in EP Publication No. 0 601 889 A2 and PCT WO 96/41166, the disclosures of which are incorporated herein.

Other complementing enzymes are known in the art, for example, pancreatic ribonuclease and Staphylococcal nuclease. Mutants of the complementing subunits of these enzymes can be constructed, by methods well-known to those of skill in the art such as site-directed mutagenesis, to generate low-affinity complementing subunits. One possible use for these types of complementing protein is as a tumor therapeutic, wherein a tumor-specific protein serves as a bridge to bring together two proteins, each of which is fused to a low-affinity complementing fragment of the nuclease. The resultant nuclease activity might, in some cases, kill the cell by destroying mRNA, genomic DNA, etc.

Binding Moieties

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Binding moieties which can be assayed for their binding affinity with each other seconds include any molecules capable of a binding interaction. The binding interaction between seconds and the seconds are seconds as a second secon the two or more binding moieties may be either direct or in the form of a complex with one the transfer of more additional binding species, such as charged ions or molecules, ligands on the transfer of the contract of tresservent exponentemacromolecules. Exponent per market environment, as the second of the second of the first second of the sec

The binding moieties which are attached to the reporter subunit can be any of a whole who was a second of the seco switch pulsars a range of different molecules including carbohydrates, lipids, proteins, and nucleic acids, as the a higher recent the well as portions, polymers and analogues thereof, provided they are capable of being linked to the reporter subunit. Exemplary proteins include members of a signal and the subunit of the subunit o transduction cascade, proteins regulating apoptosis, proteins that regulate progression of the cell-cycle or development of tumors, transcriptional regulatory proteins, translational regulatory proteins, proteins that affect cell interactions, cell adhesion molecules (CAMs), ligand-receptor pairs, proteins that participate in the folding of other proteins, and proteins involved in targeting to particular intracellular compartments, such as the Golgi apparatus, endoplasmic reticulum, ribosomes, chloroplasts and mitochondria.

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Other exemplary proteins include protein hormones and cytokines. Cytokines include those involved in signal transduction, such as interferons, chemokines, and hematopoietic growth factors. Other exemplary proteins include interleukins, lymphotoxin, transforming growth factors- α and β , and macrophage and granulocyte colony stimulating factors. Other proteins include intracellular enzymes such as protein kinases, phosphatases and synthases.

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Exemplary proteins involved in apoptosis include tumor necrosis factor (TNF), Fas ligand, interleukin- 1β converting enzyme (ICE) proteases, and TNF-related apoptosis-

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inducing ligand (TRAIL). Proteins involved in the cell cycle include deoxyribonucleic acid (DNA) polymerases, proliferating cell nuclear antigen, telomerase, cyclins, cyclin dependent kinases, tumor suppressors and phosphatases. Proteins involved in transcription and translation include ribonucleic acid (RNA) polymerases, transcription factors, enhancer-binding proteins and ribosomal proteins. Proteins involved in cellular interactions such as cell-to-cell signaling include receptor proteins, and peptide hormones or their enhancing or inhibitory mimics.

Binding of molecules will depend upon factors in solution such as pH, ionic strength, concentration of components of the assay, and temperature. In the binding assays using reporter systems described herein, the binding affinity of the binding moieties should be high enough to permit forced complementation between the reporter subunits. Nonlimiting examples of dissociation constants of the binding moieties in an assay solution, Because the state of the such as a buffered system or cell interior, are on the order of less than about 10.8 M, for the state of the second state of the state o and the second of the second section of the second of the example, less than about

15 15 10 M, or optionally, between about 10 to 10 12 M, depending upon the properties of the security is a second of particular assay system. The transport is the property of the factor of

Linking of the Reporter Subunit and the Binding Moiety

The reporter subunit and one or more binding moieties are generally linked either and are seen as the second directly or via a linker, and are generally linked by a covalent linkage. For example, when the reporter subunit and the binding moiety are proteins, they may be linked by methods known in the art for linking peptides.

In one preferred embodiment, the reporter subunit and the binding moiety comprise a fusion protein including the reporter subunit which is a low binding affinity enzyme complement and the binding moiety being assayed. The fusion protein can thus be expressed from an encoding nucleic acid intracellularly. This system is advantageous since it permits the detection and quantitation of protein-protein interactions in cells, such as mammalian cells, based on enzymatic complementation of the low affinity reporter subunits.

For example, in the embodiment wherein chimeric fused proteins are produced intracellularly that include one of two complementing low affinity β -gal mutants and a "test" protein of interest, the detected β-gal activity due to interactions between two chimeric proteins of interest will be proportional to the strength of the interaction of the

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non-β-gal protein components. Thus, the interaction is driven by the test proteins of interest, not the complementing mutants. The enzymatic activity serves as an indicator of that interaction. Another advantage of this system is that only low levels of expression of the test proteins are required to detect binding.

The fusion gene constructs preferably are constructed and transformed into cells to

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produce low level expression. The system then permits the monitoring of interactions in a given cell in the presence of endogenous competing protein partners, where the fusion protein will function as a "tracer" for the binding/association reaction. Such a system will not be prone to artifacts arising from overexpression of introduced proteins. Reduction in expression of fusion gene constructs can be accomplished by choice of appropriate promoters, ribosome binding sites and other regulatory elements. For example, fusion

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gene constructs can be introduced into vectors in which they lie upstream of an antibiotic resistance gene whose translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus internal and the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the Encephalomyocarditis virus in the land translation is regulated by the land translation is regulated by the land translation is regulated ribosome entry sequence (IRES), and which contain a mutation in the splice (1) and the sp 150 to the donor/acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the ATG sequence responsible for translational and the donor-acceptor sequences upstream of the donor-acceptor acceptor and the donor-acceptor acceptor accep initiation of the fusion gene. This type of construct results in a lower translation efficiency of the first coding sequence in a bicistronic message, but does not affect translation of the second (antibiotic resistance) sequence, which is solely dependent on the IRES. As a second (antibiotic resistance) result of these reduced levels of expression, the frequency of spontaneous interaction of

reporter subunits, which is concentration-dependent, will be significantly reduced.

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Expression of Fusion Proteins

The invention provides fusion proteins between a putative binding moiety and a low affinity reporter subunit. The putative binding moiety may comprise any protein or other molecule whose ability to bind to a second molecule is to be tested. The low affinity reporter subunit may be any molecule wherein the monomer subunit is inactive, but association of two or more identical or different monomers restores activity. The activity must be capable of generating a detectable signal. In a preferred embodiment, the low affinity reporter subunits comprise mutants of β -galactosidase capable of complementation with one another.

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Fusion proteins comprise a single continuous linear polymer of amino acids which comprise the full or partial sequence of two or more distinct proteins. The construction of fusion proteins is well-known in the art. Two or more amino acids sequences may be

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joined chemically, for instance, through the intermediacy of a crosslinking agent. In a preferred embodiment, a fusion protein is generated by expression of a fusion gene construct in a cell. A fusion gene construct comprises a single continuous linear polymer of nucleotides which encodes the full or partial sequences of two or more distinct proteins in the same uninterrupted reading frame. Fusion gene constructs generally also contain replication origins active in eucaryotic and/or procaryotic cells and one or more selectable markers encoding, for example, drug resistance. They may also contain viral packaging signals as well as transcriptional and/or translational regulatory sequences and RNA processing signals.

The fusion gene constructs of the invention are introduced into cells to assay for binding between the putative binding moieties encoded by the fusion gene constructs. The fusion gene constructs may also contain promoters and other transcriptional and/or translational regulatory sequences that are normally associated with the gene encoding the acceptance of the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that are normally associated with the gene encoding the second sequences that the second sequences is a sequence of the second sequences are sequenced to the sequences of the seque putative binding moiety. The fusion gene constructs may be introduced into cells by any method of nucleic acid transfer known in the art, including, but not limited to, viral was the same vectors, transformation, co-precipitation, electroporation, neutral or cationic liposome- and transformation, co-precipitation, electroporation, neutral or cationic liposomemediated transfer, microinjection or gene gun. Viral vectors include retroviruses, er record the poxviruses, therpesviruses, adenoviruses, and adeno-associated viruses. Particularly preferred in the present invention are retroviral vectors, which are capable of stable

integration into the genome of the host cell. For example, retroviral constructs encoding

Different fusion gene constructs encoding unique fusion proteins may be present on separate nucleic acid molecules or on the same nucleic acid molecule. Inclusion of different fusion gene constructs on the same nucleic acid molecule is advantageous, in that uptake of only a single species of nucleic acid by a cell is sufficient to introduce sequences encoding both putative binding partners into the cell. By contrast, when different fusion constructs are present on different nucleic acid molecules, both nucleic acid molecules must be taken up by a particular cell for the assay to be functional. Thus, problems of cell mosaicism are avoided when both fusion gene constructs are included on the same nucleic acid molecule.

integration and packaging signals, drug resistance markers and one or more fusion genes of

interest are useful in the practice of the invention.

The fusion gene constructs or fusion proteins of the invention may be introduced into cultured cells, animal cells in vivo, animal cells ex vivo, or any other type of cell in which it is desired to study protein-protein interactions.

Assays

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The reporter systems disclosed herein may be used to assay binding interactions of putative binding moieties attached to low affinity reporter subunits through complementation between the low affinity reporter subunits which produces a detectable signal. In addition to testing for direct binding interactions between the putative binding moieties, interactions dependent upon one or more additional molecules or ions may be evaluated. Further, multi-molecular interactions in living animal cells can be evaluated, as we are well as the influence of various drugs, peptides and pharmaceuticals on these interactions.

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In one embodiment, the binding affinity of one or more putative binding moieties स्तरिक कर कर कर may be measured by providing a reporter system including one component having one of का कार्य कर अवस्थित के 元本 と ようと かわってthe moieties bound to a low affinity reporter subunit and at least one other component との対象 かわり かんしょく かま including one other putative binding moiety bound to a second low affinity reporter and the second low affinity reporter. subunit: The binding moieties may be different or the same. In the system, the reporter of the same is the reporter of the same is the same is the system. or and degree of subunits are capable of binding and generating a detectable signal only if they are brought of the second of t The signal can assess the binding of the one or more putative binding moieties. The signal can assess the second of the be directly or indirectly detected and quantitated.

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In one embodiment of the invention, protein-protein interactions can be detected and quantitated. The signal produced by the complementing reporter subunits can serve as an indicator of binding between the putative binding moieties, either directly or indirectly via a third substance. Signals which could be detected include light emission and absorbance. Exemplary signals include chromogenic, fluorescent and luminescent signals. These signals can be detected and quantitated visually or through the use of spectrophotometers, fluorimeters, microscopes, scintillation counters or other instrumentation known in the art.

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Binding of components of the reporter systems disclosed herein will depend upon factors in solution, such as pH, ionic strength, concentration of components of the assay, and temperature. Assay solutions can be designed and developed for a particular system. The reporter systems disclosed herein can be used to conduct assays in solutions, such as buffered cell free solutions, cell interiors, solutions of cells, solutions of cell lysates, and

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solutions of cell fractions, such as nuclear fractions, cytoplasmic fractions, mitochondrial fractions, and membrane fractions. Methods for preparing assay solutions, such as enzyme assay solutions, cell extracts, and cell suspensions, known in the art may be used. For example, physiologically compatible buffers such as phosphate buffered saline may be used. See for example, the series, Methods in Enzymology, Academic Press, New York.

In one embodiment, the low affinity reporter subunits are capable of complementing one another to form an enzymatically active complex that is capable of catalyzing the conversion of a substrate to a product which is detectable, either directly or indirectly. In one embodiment, the reporter system can include two or more components, 10 each of which is a fusion protein, wherein the fusion proteins each comprise a putative binding protein fused to a low affinity reporter subunit. Thus, nucleic acids encoding the fusion proteins can be constructed, introduced into cells and expressed in cells.

அத்திரு இது அது அதிternatively, the bound reporter units or bound binding moieties can be detecting by புக்குக்குக் விறுந்தே அத the state of the bound have been specific binding moiety such as an antibody to the bound have the second and the second have 的现在分词 1.5 中国 1.

and the complementings to the second complement to the low affinity reporter subunits may be complementings to this specifies to the second complement to the second complement of the second comple subunits of β-gal. The system may include three or more reporter subunits all of which are estantes agent to the required to associate in order to produce the detectable signal. Methods for detecting the costs, as two otherwise. reaction products of active β -gal that have been developed in the art may be used. For example, β-galactosidase activity may be measured by a range of methods including livecell flow cytometry and histochemical staining with the chromogenic substrate 5-bromo-4chloro-3-indolyl β-D-galactopyranoside (X-Gal). Nolan et al., Proc. Natl. Acad. Sci, USA, 85:2603-2607 (1988); and Lojda, Z., Enzyme Histochemistry: A Laboratory Manual, Springer, Berlin, (1979), the disclosures of which are incorporated herein. Histochemical staining for β-gal can be achieved by fixation of cells followed by exposure to X-gal.

> Assays for β -gal activity described in Mohler and Blau, Proc. Natl. Acad. Sci., 93:12423-12427 (1996), the disclosure of which is hereby incorporated by reference, may be used. In one embodiment, intracellular analyses may be conducted by fixing cells and staining with the indigogenic substrate X-gal. Fixed cells also can be analyzed by assaying for β-gal activity by fluorescence histochemistry using an azo dye in combination with either X-gal or 5-bromo-6-chloro-3-indolyl β-D-galactopyranoside (5-6-X-Gal). A preferred combination is the azo dye red violet LB (Sigma Chemical, St. Louis, MO) and

5-6-X-Gal, referred to as Fluor-X-gal. For this combination, fluorescence micrographs can be obtained on a fluorescence microscope using a rhodamine/Texas Red filter set. Use of these substrates allows, for the first time, β -gal-dependent fluorescence to be visualized simultaneously with two or more other fluorescent signals.

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Vital substrates for β-gal, which can be used in living cells, are also encompassed by the invention. For example, a vital fluorogenic substrate, resorufin β-galactoside bis-aminopropyl polyethylene glycol 1900 (RGPEG) has been described. Minden (1996) BioTechniques 20(1):122-129. This compound can be delivered to cells by microinjection, electroporation or a variety of bulk-loading techniques. Once inside a cell, the substrate is unable to escape through the plasma membrane or by gap junctions. Another vital substrate that can be used in the practice of the invention is fluorescein di-β-Dgalactopyranoside (FDG), which is especially well-suited for analysis by fluorescencewww. Academics. activated cell sorting (FACS) and flow cytometry. Nolan et al. (1988) Proc. Natl. Academics and the state of the same of t

15 β-gal may also be detected using a chemiluminescence assay. For example, cells रह करिया कि अविति में containing β-gal fusions are lysed in a mixture of buffers containing Galacton Plus कि का का का का का का का का का substrate from a Galactolight Plus assay kit (Tropix, Bedford MA). Bronstein et al. (I. 1997) (1997) ணங்கள் ஊடங்கள் கிரிமார். Chemilimin., 4:99-111 (1989) the disclosure of which is incorporated herein அடி கொழிக்க நகி After addition of Light Emission Accelerator solution, luminescence is measured in a 'luminometer or a scintillation counter.

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Reporter systems other than β -gal may also be used in the practice of the invention. For example, the enzyme β -glucuronidase (GUS) can be used as a reporter and chromogenic and fluorogenic GUS substrates have been developed. The GUS substrate 5bromo-4-chloro-3-indolyl β-D-glucuronic acid (X-gluc) can be used in both chromogenic and fluorogenic applications, as follows. In one method of chromogenic staining, fixed cells are washed in PBS and stained with 2 mM X-gluc (Molecular Probes, Eugene OR), 10 mM EDTA, 0.5 mM K₃Fe(CN)₆, 0.5 mM K₄Fe(CN)₆, 0.1% Triton X-100, 0.1 M NaPO₄. Fluorogenic staining may be achieved by using a combination of 5-bromo-6chloro-3-indolyl \(\beta\)-D-glucuronic acid (5, 6 X-gluc, Molecular Probes, Eugene, OR) and Fast Red Violet LB (Sigma Chemical, St. Louis, MO). Fixed cells are rinsed with PBS and stained in 50 µg/ml 5, 6 X-gluc and 100 µg/ml Fast Red Violet LB, then rinsed in PBS.

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Fluorescence is detected on a fluorescence microscope adjusted for detection of rhodamine fluorescence.

In one embodiment of the invention, the reporter subunits comprise an enzyme and an inhibitor of the enzyme. Preferably, the inhibitor has a low affinity for the enzyme. In this case, association between the putative binding moieties is evidenced by inhibition of the activity of the enzyme. Exemplary enzymes include β-gal, GUS, β-lactamase, etc.

While dimeric reporter subunit complexes are discussed herein by way of example, multimeric reporter subunits also can be used, as can reporter subunits which are only active in the presence of one or more additional molecules or atoms. An example of a trimeric reporter subunit system would be one consisting of a β-gal ω donor (such as a ···· donor (such as a Δμ-Δω double mutant), wherein each individual mutant, and any pairwise 社会人类的特殊。 With combination of two mutants, is enzymatically inactive. Activity would be obtained only if the statements. Settle leading the first submitted and the submitted associate with one another. Enzyme reaction products can be a submitted associate with one another. Enzyme reaction products can be a submitted associate with one another. the 1800 145 and the be detected using methods available in the art, such as biochemical assay, microscopy, the state of the art, such as biochemical assay, microscopy, the state of the art, such as biochemical assay, microscopy, the state of the art, such as biochemical assay, microscopy, the state of the art, such as biochemical assay, microscopy, the state of the art, such as biochemical assay, microscopy, the state of the art, such as biochemical assay, microscopy, the state of the art, such as a state of the art, such as a state of the art, and the art, such as a state of the art, such as a state of the art, such as a state of the art, and the art, are the art, and the art, are th The methods disclosed herein enable the detection and quantitation of binding in the latest the latest the detection and quantitation of binding in the latest the la Thus, interactions between fully folded agreement as well as in intact cells. Thus, interactions between fully folded proteins are detectable, and co-translational expression of the binding moieties is not

> In the practice of the invention, the reaction product may be detected indirectly, for example, through immunological techniques, such as immunofluorescent labeling.

necessary for binding to be detected.

Protein-protein interactions can be measured in a reporter system which includes one or more fusion proteins. The fusion proteins each include a putative binding protein coupled to a low affinity reporter subunit. For intracellular expression of the fusion proteins, one or more fusion gene constructs are prepared which include sequences encoding the fusion protein(s). The fusion gene constructs may be introduced into cells by methods available in the art, including, but not limited to, viral vectors, transformation, coprecipitation, electroporation, neutral or cationic liposome-mediated transfer, microinjection or gene gun.

A variety of cell-based assays can be conducted using the cells containing the fusion gene constructs. Binding of the putative binding moieties on the fusion proteins

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expressed in the cells can be confirmed by detecting the signal produced by the reporter subunits undergoing forced complementation. Thus, for example, when the reporter subunits are complementing β-gal subunits, cells exhibiting β-gal activity indicate binding between the putative binding moieties within those cells.

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The fusion gene constructs may also contain promoters and other transcriptional and/or translational regulatory sequences that are normally associated with the gene encoding the putative binding moiety. This permits the study of physiologically-relevant · levels of the putative binding proteins in vivo, in contrast to systems in which test proteins are overexpressed. Further, this permits the study of naturally-occurring changes in levels of binding activity over time and can reveal the effects of endogenous or exogenous substances on binding interactions. Alase of the transfer of the con-

The methods and compositions of the invention can also be used to study other and the process of the molecules which influence the interaction of two putative binding partners. Proteins the confession was to Telegram described to peptides, nucleic acids, carbohydrates, lipids, ions, small molecules, synthetic compounds and a least problem 15 or the or other substances (either endogenous to the cell or exogenously added) may act as either to make the company of the cell or exogenously added). The second of a general agents are a binding interaction. By measuring the effect of such molecules of a binding interaction. By measuring the effect of such molecules on, for example, β-gal activity produced by cells containing two or more fusions and the second sec *** representing a particular pair of test proteins, agonist or antagonist activity of such a security of such molecules can be determined. Use of the methods and compositions of the invention will allow high-throughput assays to be carried out to test for agonists or antagonists of a particular binding interaction. Such high-throughput assays will be especially valuable in screening for drugs that influence medically-relevant protein-protein interactions.

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Putative binding partners, or putative binding moieties, as used in the invention, can include molecules which do not normally interact with each other, but which each interact with a third molecule so that, in the presence of the third molecule, the putative binding partners are brought together. Thus, substances which influence an interaction between putative binding partners include those which stimulate a weak interaction between putative binding partners, as well as one or more molecules which mediate interaction between molecules which do not normally interact with each other. In addition, substances which influence an interaction between putative binding partners can include those which directly or indirectly affect an upstream event which results in association between the putative binding partners. For example, if phosphorylation of one of the

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putative binding partners endows it with the capacity to associate with another of the putative binding partners; substances which influence the interaction of the putative binding partners include those which directly or indirectly affect a kinase activity.

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Assays can be developed as disclosed herein to examine the effect on intermolecular interactions of a variety of compositions including drugs such as antipyretic and anti-inflammatory drugs, analgesics, antiarthritics, antispasmodics, antidepressants, antipsychotics, tranquilizers, antianxiety drugs, narcotic antagonists, antiparkinsonism agents, cholinergic antagonists, chemotherapeutic agents, immunosuppressive agents, antiviral agents, parasiticides, appetite suppressants, antiemetics, antihistamines, antimigraine agents, coronary vasodilators, cerebral vasodilators, peripheral vasodilators. hormonal agents, contraceptives, antithrombotic agents, diuretics, antihypertensive agents, cardiovascular drugs, opioids, and vitamins. • AND SERVICE CONTRACTOR

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non-discretive the sale-sket of the Protein-protein interactions mediated by a third molecule can be detected and the sale and the second and th and 2 below, wherein β-gal fusion proteins are used to monitor of 146 and 2 below, wherein β-gal fusion proteins are used to monitor of 146 and 2 below, wherein β-gal fusion proteins are used to monitor of 146 and 2 below. and the rapamycin-mediated interaction of the FKBP12 and FRAP proteins. Belshaw, P. J. et also and a second proteins. 201 April 1996 | Al., Proc. Natl. Acad. Sci. USA, 93: 4604-4607 (1996); Brown et al., Nature 369: 756-758 Company of the control of the contro Science, 273:239-242 (1996). For example, kinetics of binding can be determined by 20 measuring β -gal activity at different times following addition of rapamycin to cultures of cells expressing fusions of FKBP12 and FRAP to two complementing, low affinity β-gal mutants (e.g., $\Delta \alpha$ and $\Delta \omega$). A dose-response curve can also be obtained, in which the extent of binding, as measured by β-gal activity, is determined as a function of rapamycin concentration.

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This assay can be adapted to control for the possible effect of a protein component on its fusion partner, thereby enabling the study of protein-protein interactions in a quantitative fashion. In one such control system, tripartite fusion constructs including a reporter subunit, a binding protein and the protein of interest are provided. As described below in Example 3, in one embodiment, the fusion protein includes 1) a β-gal mutant portion, 2) a FKBP12 or FRAP portion, and 3) a test protein portion. The most N-terminal component is the test protein, followed by FKBP12-Δω or FRAP-Δα. The presence of FKBP12 and FRAP in these constructs allows rapamycin-mediated dimerization of the

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fusion proteins. The absolute values of β-gal activity obtained by simple co-expression of a fusion containing a test protein of interest and fusions containing different potential interacting partners is determined. In parallel samples, β-gal activity is measured upon induction of complementation with a fixed amount of rapamycin. The ratio of \beta-gal. activity obtained in the absence and the presence of rapamycin indicates the relative abilities of the different protein pairs to interact with each other.

A further advantage of the tripartite fusion system is that the presence of the FKBP12 and FRAP components provides a flexible hinge domain between the β-gal mutants and the test protein. This reduces the possibility of interference between the B-gal accomponent and the test protein. Furthermore, it allows direct testing of the functional integrity of the β-gal components in the fusions without the need for recloning into more. A second efficient viral vectors. For example, the tetracycline repressor, tetR, forms homodimers in the state of the அத்து சென்ன சென்னைகள்கால் ells with high-efficiency. Hinrichs et als (1994) Science 264:418-420 இரும் வெள்ள அதிக்கு வருக்கு parameters and Coexpression of tetR-FKBP12-Δω and tetR-FRAP-Δα fusions yielded β-gal-positive cells, which is the second of tetR-FKBP12-Δω and tetR-FRAP-Δα fusions yielded β-gal-positive cells, which is the second of tetR-FKBP12-Δω and tetR-FRAP-Δα fusions yielded β-gal-positive cells, which is the second of tetR-FKBP12-Δω and tetR-FRAP-Δα fusions yielded β-gal-positive cells, which is the second of tetR-FKBP12-Δω and tetR-FRAP-Δα fusions yielded β-gal-positive cells, which is the second of tetR-FKBP12-Δω and tetR-FRAP-Δα fusions yielded β-gal-positive cells. Assessment of the State of the N-terminal peptide component efficiently drives (1967) complementation of the C-terminal mutant β-gal polypeptides, with the FKBP12 and

The system may be further tested and compared by constructing fusions between Programme and the second of the second each β-gal mutant and the complete coding sequence of MEF2c. Since MEF2c is known to form homodimers in vivo, coexpression of both β-gal mutants fused to MEF2c should result in readily detectable enzymatic activity. MEF2c mutants that are impaired in their dimerization potential are available and fusion of one of the mutants to one of the \beta-gal' mutants can serve as a negative control to further validate the system. Molkentin, et al., Mol. Cell. Biol., 16:2627-2636 (1996).

> The reporter system can also be designed with controls to permit the quantitation of the expression level of the β-gal fusion proteins. This will make it possible to control for potential differential expression of the two (or more) fusion proteins. For example, a peptide tag for which well-characterized monoclonal antibodies are available may be fused in frame at the C-terminus of each \beta-gal mutant. Different tags, such as flag and myc may be used for $\Delta\alpha$ and $\Delta\omega$, to allow differential detection of the two mutants even when coexpressed in the same cells. In parallel with the determination of β -gal activity in the

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lysates of these cells, an ELISA assay can determine the precise amount of each \(\beta \)-gal fusion protein in the same lysates. First, a polyclonal anti-β-gal antiserum may be used to immobilize the antigens. Then the monoclonal antibody directed against the appropriate tag followed by an enzyme-linked anti-mouse secondary antibody may be used to quantify the amount of the β-gal fusion protein of interest. Such an approach, employing wellcharacterized techniques, should allow a determination of the expression levels of each fusion protein. This modification will be useful where the attached tag does not impair the binding of the protein or the ability of the reporter subunits to complement.

Applications of the Invention

As will be apparent to one of skill in the art, the invention allows, for the first time, a broad range of studies of multiprotein and other types of multi-molecular interaction to was a street of the be carried out quantitatively or qualitatively in living cells. In what follows, non-limiting: அத்துக்கு கார்க்கு examples of different applications of the invention are provided. வகுக்கு கொள்ள அத்துக்கு காக்குக்கு காக்கு காக

. 1948 Washington 12 Percention that levels of β-gal activity in the presence and absence of forced department about the control of the cont 15. 15. 1 to complementation can be distinguished, both biochemically (Figure 5) and by FACS To be a local and the second Example 10 and Figure 6), suggests that the methods of the invention can be used to a second construction of the invention can be used to a second construction of the invention screen for new binding partner(s) for a given target protein. In this embodiment, the target characterized cell line. Expression libraries containing cDNAs fused to a weaklycomplementing β-gal mutant are introduced into these cells using, for example, retroviral vectors (e.g., Kitamura et al., Proc Natl. Acad. Sci. USA 92:9146-9150 (1995)) or any other means of gene transfer known in the art. Vectors expressing gene products that interact with the target protein are isolated by identifying β-gal-positive clones. An advantage of this system is that the screen can be carried out in any cell type, regardless of the cell's milieu of endogenous (and potentially competing) proteins. A further possibility for this type of system is that the target protein can be localized to a specific cellular compartment, with the aim of identifying proteins involved in interactions restricted to that particular location.

> The use of fluorescence-activated cell sorting techniques is particularly well-suited to this embodiment of the invention. For example, β -gal-positive cells which contain cDNAs expressing gene products that interact with the target protein will generate a signal that will allow such cells to be purified by cell-sorting techniques. Such cDNAs could be

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delivered, for example, using retroviral vectors that allow introduction of high complexity cDNA libraries with high infection efficiency.

The assays and methods of the invention can also be carried out in the presence of extracellular signaling molecules, growth factors or differentiation factors, peptides, drugs or synthetic analogs, or the like, whose presence or effects might alter the potential for interaction between two or more given proteins in a particular cell type.

Detection of molecular interactions, using the methods and compositions of the invention, is not limited to those occurring in the nucleus, nor is it limited to intracellular interactions. For instance, interactions involving surface receptors can be detected in the 10.000 as practice of the invention. In one embodiment, the invention provides new techniques for a second edetecting ligand-induced dimerization of surface receptors in living cells. Dimerization, or and the surface receptors in living cells. higher order oligomerization, of cell surface receptors is often a prerequisite for receptor agreements to the contraction and ensuing signal transduction. (For example, the binding of epidermal growth has the requirements of the contraction and ensuing signal transduction. அது நக்கு கொண்டுக்கு factor (EGF) to its receptor stabilizes the dimerization of the receptor and leads to ach அக்கிய கொண்டுக்க கொண்டுக்க கொண்டுக்கு முற served and 15 meter activation of its tyrosine kinase activity. Schlessinger et al. (1992) Neuron 9:383-391; and served activity and served activity. and Weiss et al. (1991). Gell 61:203-212; and Weiss et al. (1997) Curr. Opin. Genet. Devices algered in the lines. 7:80-86. Example 11, infra, discloses the use of β-gal complementation to monitors waster the weakly a membrane receptor dimerization in living cells. For this purpose; the weakly a construction of the large and a construction of the living cells. complementing $\Delta\alpha$ and $\Delta\omega$ deletion mutants of β -gal were fused to the extracellular and α transmembrane regions of the human EGF receptor to form a chimeric receptor molecule (see Figure 7A). Deletion of the cytoplasmic domain of the receptor prevents the internalization and degradation of the receptor that is normally observed following EGF stimulation of cells (Livneh et al. (1986) J. Biol. Chem. 261:12490-12497), permitting an analysis of receptor dimerization over time in changing conditions. The results presented in Example 11 demonstrate that this embodiment of the invention can be used to detect a previously-unrecognized mode of regulation of EGF receptor signaling, in which EGF receptor tyrosine kinase activity acts as a feedback inhibitor limiting the dimerization of the receptor.

The practice of the invention is not limited to detection of interaction between two different molecules. Multimerization of a molecule can also be detected using the methods and compositions of the invention. In this regard, Example 11 discloses the detection of receptor dimerization through the practice of the invention.

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By combining the methods and compositions of the invention with state-of-the-art methods for construction of high-titer, high-complexity cDNA libraries in retroviruses (e.g., Pear et al., (1993) Proc. Natl. Acad. Sci. USA 90:8392-8396), it will be possible to identify interaction partners of a specific test protein in mammalian cells (i.e., perform functional genomics at the protein level). For this application, construction of cDNA libraries in retroviral vectors wherein the cDNA coding sequence is fused to a sequence encoding a low affinity reporter subunit will be used. A sequence encoding a binding protein of interest will be fused to a low affinity reporter subunit in a first retroviral vector. In a second series of retroviral vectors, a second complementing low affinity reporter 10 subunit will be fused to a variety of different proteins that will be tested for their ability to bind to the protein of interest. Testing will be conducted by co-infection of cells with the first and one of the series of second retroviral vectors. Those test proteins which are Apply and the protein of interest will allow detection of a reporter signal in cells one to be a considerable of binding to the protein of interest will allow detection of a reporter signal in cells one to be a considerable of binding to the protein of interest will allow detection of a reporter signal in cells one to be a considerable of binding to the protein of interest will allow detection of a reporter signal in cells one to be a considerable of binding to the protein of interest will allow detection of a reporter signal in cells one to be a considerable of binding to the protein of a considerable of binding to the protein of a considerable of binding to the protein of a considerable of the c Research and a specific which they are co-expressed with the protein of interest. This application will also be an in the acceptance of the control of the c で policy 15点。 Spruseful in screening for agonists and antagonists of medically-relevant protein interactions はみない。 多の こうかい with the second of the invention, cells in which a protein encoded by one of the agent as the many THE COURT OF SERIES OF SECOND VECTORS is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of second vectors is able to interact with the binding protein of interest encoded by the series of the second vectors is able to interact with the binding protein of interest encoded by the second vectors is able to interest encoded by the second vectors is able to be agreed the control of the first vector are detected and isolated by flow cytometry or fluorescence-activated cells and account of the control sorting (FACS). Methods for flow cytometry and FACS are well-known in the art; e.g., Nolan et al. (1988) Proc. Natl. Acad. Sci. USA 85:2603-2607; Webster et al., Exp. Cell Research, 174:252-265 (1988); and Parks et al. (1986) in The Handbook of Experimental Immunology, (eds. Weir, D.M., Herzenberg, L.A., Blackwell, C.C. & Herzenberg, L.A.), Blackwell, Edinburgh, 4th edition, pp. 29.1-29.21. In this way, clones of cells in which binding occurs can be isolated and propagated for further study. This aspect is particularly suited for studies of developmental mechanisms, wherein it is possible to select a population of cells in which a particular developmentally-relevant interaction has occurred and study the further development of that cell population, while at the same time, studying the further development of cells in which the interaction has not occurred, for comparison. In a similar fashion, the practice of the invention makes it possible to isolate and/or study the further development of cells exhibiting interactions involving protein such as transcriptional regulatory proteins, translational regulatory proteins, DNA replication proteins, mRNA splicing proteins, proteins involved in signal transduction, proteins

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involved in cell-cell and cell-substrate adhesion (for example, cell movement, axon guidance and angiogenesis), oncogene products, tumor suppressors, proteins involved in cell-cycle control and viral proteins, such as those involved in regulation of viral replication, virus-host interactions and virus assembly, and proteins which are subunits, crosslinkers, modifying agents or molecular motors within the cytoskeleton of cells.

For a given target protein whose gene is capable of being fused to a low-affinity complementing reporter subunit, it is possible to identify known and heretofore unknown proteins or other endogenous or extraneous substances with which it interacts, by using the compositions and methods of the invention. In like manner, for a sequence which encodes a protein of unknown function, such as may be obtained from a nucleic acid sequence database, (or a plurality of sequences such as a cDNA library) the practice of the invention allows one to identify molecules with which the encoded protein interacts. The identity of குறும் கடிக்கத்திக்கு the interacting molecule(s) is likely to provide information with respect to the structure மக்கத்திக்கு சிக்க as leaves or about the practice of the invention will likely aid where a respective of the invention will likely aid where a respective of and agree 15 may report in the identification and characterization of newly-discovered proteins and protein-coding at the gipes in American a representative of nucleic acid sequences. Facility was a respect to the control of the control

कार कार्या के किया है जिस्सा another aspect of the invention, a shotgun approach to the identification of कार्या के किया कार्या करते. research responsible or protein-protein interactions can be taken by generating a first set; of constructs which will research and research express the encoded products of one cDNA library fused to a first low-affinity complementing subunit and a second set of constructs which will express the encoded products of a second (or the same) cDNA library, fused to a second low-affinity complementing subunit. Co-expression of the two sets of constructs and selection of cells in which complementation occurs will allow the isolation of clones and the identification of cDNAs which encode interacting partners. One or both of the interacting partners may. be known; alternatively, both of the interacting partners may represent heretofore unidentified proteins. If both partners are known, new information about their binding specificity may be obtained. If one partner is known, it may provide information on the function of the unknown binding partner. If neither are known, the observation that they interact may assist in the eventual identification of one or both of the interacting pair.

The invention may be applied to studies of the mechanisms that regulate either homo- or hetero-dimerization or multimerization of specific molecules, including high

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efficiency screening to identify synthetic or naturally occurring compounds capable of influencing such dimerization.

The invention can be used for investigations relating to the localization of specific complexes within intact cells, or intact animals. Types of cells which can be used are primary or established cell lines and other types of embryonic, neonatal or adult cells, or transformed cells (for example, spontaneously- or virally-transformed). These include, but are not limited to fibroblasts, macrophages, myoblasts, osteoclasts, osteoclasts, hematopoietic cells, neurons, glial cells, primary B- and T-cells, B- and T-cell lines, the state of the chondrocytes, keratinocytes, adipocytes and hepatocytes.

It is also possible, through practice of the invention, to devise systems for regulation of enzyme activity by regulating the association of complementing mutants. This aspect of the invention has potential applications to human therapy, as a method to applications to human therapy, as a method to அது புக்கத்தில் கொள்ள Cregulate the enzyme-driven conversion of pro-drugs into their active forms? இதுக்கு மக்க கொள்ள கொள்ள அதாக்குள்ளது. கூடு, அதி நக்கு குறை Processes involving molecular interactions, particularly protein-protein குறைக்கு கொண்டு கூடுக்கு இது விக்கு Hardword 15 end was interactions, which can be studied in the practice of the invention include, but are notice. The practice of the invention include, but are notice. The practice of the invention include, but are notice. particle was self-limited to, transcription, translation, replication, mitosis, growth control, progression and transcription, translation, replication, mitosis, growth control, progression and translation and t regulation of the cell-cycle, apoptosis, cell-cell, cell-substratum and cell-ligand এ স্থোক্তিক ক্রম্প্রাnteractions, intracellular signal transduction cascades, oncogenesis, cell·lineages, and নি ক্রমিণ্ডেই প্রকর্তনার করিব embryonic development. Examples of cell ligands include leptin and growth factors such as epidermal growth factor (EGF), nerve growth factor (NGF), platelet-derived growth factor (PDGF), and insulin-like growth factors I and II (IGF-I and IGF-II), transforming α growth factors α and β (TGF- α and TGF- β), endorphins and endorphin receptors, prostaglandins and their receptors, cytokines and their receptors, neurotransmitters and their receptors, adrenergic receptors, and cholinergic receptors. Receptors which could interact with ligands include EGF, NGF, and PDGF receptors and leptin receptors. Analysis of EGF receptor dimerization, using the methods and compositions of the invention, is provided in Example 11, infra.

Additional interactions that can be studied by the practice of the invention include interactions involved in cell metabolism and cell structure. These include, but are not limited to, interactions that are involved in energy metabolism or which establish or modify the structure of the membranes, cytoplasm, cytoskeleton, organelles, nuclei, nuclear matrix or chromosomes of cells. Interactions among constituents of the

extracellular matrix, or between constituents of the extracellular matrix and cells, can also be studied with the methods and compositions of the invention.

The invention will be further understood by the following non-limiting examples.

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EXAMPLES

Example 1: Preparation and Transfection of Retroviral Construct Encoding a β-Galactosidase Reporter System.

A reporter system using β-galactosidase ("β-gal") complementation to evaluate protein-protein interactions was constructed. A well-characterized protein complex developed by Schreiber was used as a test system to provide the protein binding moieties. Belshaw, P. J. et al., Proc. Natl. Acad. Sci. USA, 93: 4604-4607 (1996); Brown et al., Acad. Sci. USA, 93: 4604-4607 (1996); Brown et al., Nature 369: 756-758 (1994); Chen, et al., Proc. Natl. Acad. Sci., USA, 92:4947-4951 (1995); and Choi, J. et al., Science; 273:239-242 (1996); the disclosures of which are முது நார்கள் கண்ணார் நார்கள் incorporated herein. In this protein complex, the intracellular rapamycin binding protein, கூடுக்கு செரு இதிரும் 15. 15. 15. FK506-binding protein-12 (FKBP12), interacts with intracellular FKBP-rapamycin in the control of th associated protein (FRAP) only when rapamycin is present in the culture medium, an Acade Sand Control of Sand interaction that increases with the dose of rapamycin. Rapamycin is a small, cell-permeable special residence of the two intracellular proteins via independent determinants. Since the two intracellular proteins via independent determinants. Since the two intracellular proteins via independent determinants. rapamycin is unable to bind two FKBP12 molecules at the same time and FRAP only binds rapamycin within the FKBP12-rapamycin complex, only heterodimers should form upon rapamycin treatment. Ho, S. N. et al., Nature, 382:822-826 (1996), the disclosure of which is incorporated herein.

> The β-gal system was combined with the FKBP12/FRAP/rapamycin system as follows. Two different retroviral constructs were produced, each encoding fusion proteins of the $\Delta\omega$ or $\Delta\alpha$ β -gal mutants, and either FKBP12 or the FKBP-rapamycin binding domain of FRAP, respectively (FKBP12- $\Delta\omega$ and FRAP- $\Delta\alpha$).

The $\Delta\alpha$ or $\Delta\omega$ β -gal mutants were obtained as described in Mohler and Blau, *Proc.* Natl. Acad. Sci., 93:12423-12427 (1996), the disclosure of which is incorporated herein.

To fuse the sequences coding for FKBP12 and the FKBP12-rapamycin binding domain in frame with β-gal, an adapter oligonucleotide (CATGGAGCTCGAGAG) containing an XhoI site was inserted in the NcoI site at the ATG of the Δα and Δω β-gal mutants described by Mohler and Blau, supra. Two XhoI-SalI DNA fragments

corresponding to amino acids 2025-2114 of human FRAP and to the complete coding sequence of human FKBP12 were cloned in the XhoI site of the modified $\Delta\alpha$ and $\Delta\omega$ mutants, generating FRAP- $\Delta\alpha$ and FKBP12- $\Delta\omega$. Conservation of the appropriate reading frame was confirmed by sequencing for both constructs.

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To insert the FRAP-Δα and FKBP12-Δω coding sequences in the pWZL-Neo and pWZL-Hygro retroviruses, an adapter oligonucleotide containing NcoI and BamHI sites (GATCACCATGGACGCGTGGATCCC) was inserted in the BamHI and XhoI sites of the pWZL vectors. Both the original sites were destroyed by this insertion. The FRAP- $\Delta\alpha$ and FKBP12-Δω coding sequences were then inserted in the modified pWZL vectors as Burgard Burgard State Control NcoI-BamHI fragments.

The cDNAs encoding FKBP12-Δω and FRAP-Δα were inserted into a mouse ecotropic retroviral vector upstream of the hygromycin resistance or neomycin resistance As yours and the surgences, respectively, as described above. By using an Encephalomyocarditis virus internal on the appropriate fraction. প্ৰায়ে লোগৰ কৰি ribosomal entry sequence (IRES), introduction of a single retroviral vector ensured নি কৰিব কৰিব কৰিব কৰিব ा production of a bicistronic mRNA and translation of both the Δα –β-gal-FRAP-protein and ्राप्ति अस्ति अस्तिक # 10 March 10 Control of the drug selectable hygromycin protein. A second retroviral vector yielded the Δω-β-gal- () in the land of the AND FKBP12 protein and neomyciń resistance protein.

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প্ৰস্তুত্বিপ্ৰত কৰে ক'বছ ক'বছ For virus production and infection, provinal constructs were introduced into হৈ সুকলি বুলি বুলি বুলি কিন্তুত্বি packaging cells by calcium phosphate transfection. The supernatant media containing retrovirus from the packaging cells was harvested 24 to 72 hours after transfection and used to infect C2C12 cells in the presence of 8 µg/mL polybrene. Singly and doubly infected cells were selected with the appropriate drugs. Both Geneticin and Hygromycin were used at a final concentration of 1 mg/ml. The selected cells were expanded as populations for subsequent experiments.

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Although the background β -gal detected with the $\Delta\alpha$ and $\Delta\omega$ mutants expressed from MFG retroviral vectors described previously (Dhawan et al., Science, 254:1509-1512 (1991) was relatively low (Mohler, W. A., & Blau, H. M., Proc. Natl. Acad. Sci. USA, 93:12423-12427 (1996), the disclosure of which is incorporated herein), it was further reduced by using retroviral vectors with point mutations that deleted the splice donor/acceptor sequences upstream of the β -gal ATG (pWZL). These mutations result in a lower translation efficiency of the first coding sequence contained in the vector, but do not affect the expression of the selectable marker, which is solely dependent on the IRES.

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Using this vector, two-fold less of the upstream protein was expressed compared to vectors containing the same LTRs (long terminal repeats) and the wild-type splice donor/acceptor sequences. Such a reduction in translation reduces the concentration of the fusion protein and consequent spontaneous interactions of β-gal mutants irrespective of the test proteins to which they are fused. As a result, in preliminary experiments, the background enzyme activity measured by luminometer for $\Delta\alpha$ and $\Delta\omega$ β -gal mutants alone was reduced from low to essentially undetectable.

Infectious viral particles were produced by transient transfection of each construct shown in Figure 2a into a packaging cell line modified from that described by Pear.et al., (1993) Proc. Natl. Acad. Sci. USA 90:8392-8396 by calcium phosphate transfection. The supernatant media containing retrovirus from the packaging cells was harvested 24 to 72 hours after transfection and used to infect C2C12 cells in the presence of 8 μg/mL γ μγ (λομά (ed newer series polybrene. C2C12 myoblasts were infected either singly with each retrovirus alone or a series and a series of the series of th Assess and the selection with all experiments were performed after selection with all assessments the assessments are a selection with all assessments the assessments are a selection with a sel 15 15 how hygromycin and G418 to ensure that 100% of the cells contained both constructs. Both 100% of the cells contained both constructs. Both A transfer of the selected of cells were expanded as populations for subsequent experiments. A comparation of the compa

Example 2: Assays of Binding and Activity of the β-Galactosidase Reporter of the β-Galactosid System.

Following the addition of rapamycin to the media, the transfected cells obtained as described in Example 1 were assayed for β -gal activity. As shown in Figure 3, C2C12cells expressing both FKBP12-Δω and FRAP-Δα were tested by exposure to 10 ng/ml rapamycin (Figure 3b) for 12 hr or to no drug at all (Figure 3a). Only those cells expressing both constructs exhibited \beta-gal activity, readily visualized by X-gal staining of fixed cells (Figure 3b). It is advantageous that cytoplasmic staining is detectable with this method, in contrast to prior methods such as the yeast two-hybrid system, which report only nuclear interactions. X-gal staining was conducted as follows: Cells were fixed 5 minutes in PBS plus 4% paraformaldehyde and rinsed in PBS prior to staining. Indigogenic X-gal staining was performed overnight at 37°C in PBS plus 1 mg/mL X-gal, 1 mM MgCl₂, 5 mM K₃Fe(CN)₆, 5 mM K₄Fe(CN)₆.

The kinetics of activation of β-gal upon rapamycin treatment were determined. C2C12 cells expressing both fusion proteins were plated in replicate in 96 well plates.

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Rapamycin was added to the culture medium, and the β-gal activity measured at different time points. For each time point, six replicate samples were assayed with a sensitive chemiluminescence assay, as described in Mohler, W. A., & Blau, H. M., Proc. Natl. Acad. Sci., USA, 93:12423-12427 (1996), the disclosure of which is incorporated herein. In the assay, cells cultured in microtiter plates were lysed in situ in 50 µL of a 1:3 mixture of lysis and assay buffers containing Galacton Plus substrate from the Galactolight Plus assay kit (Tropix, Bedford, MA). Reactions proceeded for 1 hour at room temperature. - After addition of Light Emission Accelerator solution, luminescence was measured in a scintillation counter.

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10. The results, shown in Figure 4, indicate that the interaction assays for the fusion proteins are specific, and exhibit similar kinetics and a comparable dose-response curve to ്ളെ ുപ്പാരുക്ക് shours. As a control, no rapamycin was added, and no β-gal·activity was detected above കാര്യ്യ് അവരുക്കും and the state of t ு சாவர்க்கு அக்கு அக்களை ructs, β-gal activity did not increase above background when rapamycin was added கூண்ணுக்க கணையாக

In Figure 4b, the dose response curve is shown. The activation of β-gal was worth a street of the s which appeared linear over a range of 0 to 10 ng/ml of the drug. This linearity provides support that β-gal enzymatic activity can serve as a reporter to quantitate protein-protein interactions. The close approximation of both the dose response and the kinetics to that observed by others (Ho, S. N. et al., Nature, 382:822-826 (1996)) suggests that the fusion to β -gal peptides is not interfering with the interaction of the FKBP12 and FRAP proteins. Moreover, endogenous FKBP12 and FRAP proteins are ubiquitously expressed and are capable of interacting with each other or with the fusion proteins in the presence of rapamycin, without generating β -gal activity. Detection of β-gal activity, as shown above, indicates that productive FRAP-Δα and FKBP12-Δω dimers will form in a cellular environment containing competing endogenous proteins, and that the resultant β-gal activity reflects the interaction of FRAP and FKBP12rapamycin Thus, the β-gal fusion proteins can be used to monitor the interaction of proteins in the FKBP12/FRAP/rapamycin complex and in other types of multiprotein complexes.

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It is also possible to detect and quantitate binding activity in cell lysates. As shown in Figure 5, cells expressing both FKBP12-Δω and FRAP-Δα fusion proteins were expanded in the absence of rapamycin and lysed. 100 ng/ml rapamycin was added to half of the samples, and the β-gal activity in the treated and untreated lysates was determined immediately (white bar), after one hour (black bar) or after 3 hours (gray bar). A greater than two-fold increase in β-gal activity was observed in the rapamycin-treated lysates one hour after administration of the drug. In control lysates that were not exposed to rapamycin, no statistically significant increase in β-gal activity was detected. The ability to detect and quantitate protein-protein interactions in cell lysates using the methods and compositions of the invention indicates that interactions between mature, fully-folded proteins can be detected and quantitated; co-translational assembly of complexes in not required for formation of complexes that can be monitored by β-gal activity.

Example 3: Tripartite fusions for the quantitation of protein-protein interactions of the protein interaction in the protein interaction of the protein interaction in the protein in the pro

तकार विकास का का का का कि permit protein interactions to be studied in a quantitative manner in the system स्थान अवस्थित अवस्थान । ംൻ വരു അംഗ പ്രീട്ട കുറിയുടെ adescribed in the above Examples and to control for effects on either the binding ability of ് ആ ുക്കുന്നു വരുടെ പ്ര தாகை முத்துக்குக்க the binding moiety.or the complementing ability of the reporter subunits resulting from கண்ணன் செய்ய நடி The particular both activities being present in a single fusion/protein, additional modifications were made Approximately and the particular and t as the system of the above described system with the above described system with the above described system with the system of the system. the β-gal fusion proteins will be expressed from the same viral promoter, however, for the same viral promoter, however, however, for the same viral promoter, however, ho some proteins, it is possible that their expression level will be influenced by the specific fusion partner. In particular, some proteins or domains could affect the stability or conformation of the β-gal domain. As a result, differences in the ability of the test proteins (the putative binding moieties) to complement one another could be observed that are not based on a physiological mechanism.

> In order to avoid these problems, fusions containing three components (β-gal mutant, FKBP12 or FRAP, and the test protein) were constructed. The most N-terminal component is the test protein, followed by FKBP12- $\Delta\omega$ or FRAP- $\Delta\alpha$ (see the exemplary system in Figure 2b, where the test protein portions of the fusion are indicated by X and X'). The presence of the FKBP12 and FRAP portions allows rapamycin-mediated dimerization of these fusions, and the efficiency of β -gal complementation in the presence of rapamycin appears to be dependent on the FKBP12/FRAP/rapamycin interaction. The absolute values of β-gal activity obtained by simple coexpression (in the absence of

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rapamycin) of fusions containing a fixed protein of interest and different interacting partners was determined. In parallel samples, β-gal activity was measured upon induction of complementation with a fixed amount of rapamycin. The ratio between the β -gal activity obtained in the absence or in the presence of rapamycin indicated the relative ability of the different protein pairs to interact with each other. An added advantage of this approach is that the presence of the FKBP12 and FRAP domains provide a flexible hinge between the β -gal mutants and the putative binding moieties that are being analyzed. This reduces the possibility of interference between β -gal and the proteins of interest. Furthermore, it allows direct testing of the functional integrity of the β -gal components in the fusions without the need for recloning into more efficient viral vectors. A DO NOTE OF THE SEC.

Results were obtained with tetR-FKBP12-Δω or tetR-FRAP-Δα tripartite fusions & complete to the second seco (see example in Figure 2b). Coexpression of these constructs, in which dimerization is see the constructs of the constructs. exercises represented 420 (1994); the disclosure of which is incorporated herein); readily yielded β-gal positive of a negative for the disclosure of which is incorporated herein); 15. 2 macells: This indicates that functional tripartite fusions can be constructed, in which the Adaptive of the constructed in and the first of the sent sent that the sent of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficiently drives complementation of the most N-terminal peptide component efficient சு சார்ப் கூறி நடித்து and the C-terminal β-gal-deletion mutant polypeptides. இது நடித்து நடித்து

ない。 ないは、 できない また ない Example 4: <u>Dimerization of myogenic regulators using complementing 段 gal かけっぱい ないできません</u> fusion proteins

The β -gal complementation system is used to assay for the dimerization and nuclear translocation of HLH proteins (helix-loop-helix proteins, Murre et al. (1989) Cell 56:777-783) including activators of muscle-specific proteins (myoD, myogenin, myf5, MRF-4), inhibitors of myogenesis (Id, Mtwist, I-mf) and ubiquitous E2A-type proteins (E47, E12, HEB).

In a first step, a myoD-Δα-β-gal (myoD-Δα) fusion construct and a E12-Δω-β-gal $(E12-\Delta\omega)$ fusion construct are engineered in selectable retroviral vectors, as described above for FRAP- $\Delta\alpha$ and FKBP12- $\Delta\omega$. The two constructs are transduced into C2C12 myoblasts. Following selection with the appropriate drugs for cells which express both constructs, β -gal activity is quantitated using the chemiluminescent assay described above. β-gal activity indicates that heterodimerization of the fusion proteins is occurring in this cell type. If \beta-gal activity is detected, individual cells are analyzed using a fluorescent Xgal stain in order to determine if the heterodimers are present in the nucleus. Since wild-

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type β-gal can be specifically directed to and detected in the nucleus by inclusion of a nuclear localization sequence (nls) (Hughes and Blau, Nature, 345:350-352 (1990)), activity from the β -gal hybrid protein may be detected in the nucleus. Knowledge of the site of localization in the cytoplasm or nucleus will aid in assessing the function of the protein interactions, e.g. sequestration and inhibiting activity, or promoting activity. This method permits visualization of fluorescent markers of myogenesis, such as desmin, and creatine kinase, in correlation with the localization of β -gal, using the sensitive Fluor-X-Gal substrate described above (Mohler, W. A., & Blau, H. M., Proc. Natl. Acad. Sci., USA, 93:12423-12427 (1996)). 11.44 (12.14)

10 All fusion constructs between myogenic regulators and complementing β -gal . We mutants described in the following sections may be tested in a muscle cell where we have the section of the following sections are the section of the following sections are the section of the section of the following sections are the section of the section heterodimerization of the endogenous myogenic regulator is known to occur. In addition, A state of the following controls also may be performed. The myoD-Δα construct may be the following controls also may be performed. The myoD-Δα construct may be the following controls also may be performed. 14.00% of the season contransduced into the cell with FKBP12-Δω, and the E12-Δω construct may be it wise an expension of the cell with FKBP12-Δω, and the E12-Δω construct may be it wise a season of the cell with FKBP12-Δω, and the E12-Δω construct may be it wise a season of the cell with FKBP12-Δω, and the E12-Δω construct may be it wise a season of the cell with FKBP12-Δω, and the E12-Δω construct may be it wise. ு நாருக்கும் விருந்து cotransduced with FRAP-Δα. This combination of constructs should result in no β-gal கொண்ட கிறு சிறு அதி ந The second process activity, unless some unusual mechanism exists in the particular cell type being tested that the control of the second process activity. enhances complementation of the weakly complementing β-gal peptides independent of 下了了一个文章的 heterodimerization of the non-β-gal parts of the molecule. The FRAP-Δα and FKBP12-1 (東京中央市場大阪市場 $\Delta\omega$ may also be cotransduced and cells treated with rapamycin as a positive control for complementation in each cell type. Cells in high serum medium (growth medium) and cells in low serum medium (differentiation medium) should/will give different results.

> Example 5: In vivo assay for the effect of growth factors and substrates on heterodimerization and homodimerization.

Using the constructs described above in Example 4, C2C12 myoblasts are transduced with one of the myogenic HLH fusion constructs and the E12-Δω construct. Although C2C12 cells will already contain endogenous myogenic HLH proteins and E12, the chimeric constructs will act as a "tracer" to measure the extent of heterodimerization. Transduced cells then may be stimulated to either differentiate or proliferate by changes in serum levels or the addition of growth factors (TGF-\beta, bFGF, IGF-I and IGF-II) in the presence or absence of substrates such as fibronectin or laminin. β-gal activity then is measured as a function of time. Rapid changes in β-gal activity after growth factor stimulation may suggest a more direct mechanism of action of a given extracellular signal

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on the formation of specific heterodimers. Slower changes may indicate that the extracellular signal acts indirectly, for example by up-regulating the expression of a competing factor which can sequester one or both fusion proteins. Changes in β-gal activity may be correlated with the expression levels of known inhibitors of differentiation such as Id proteins, measured by Northern blot in parallel samples. A comparison of the kinetics of changes in β-gal activity obtained with each pair of test proteins in parallel experiments will indicate whether specific MRFs (muscle regulatory factors, Yun et al. (1996) Curr. Opin. Cell Biol. 8:877-879; and Cossu et al. (1996) Trends Genet., 12:218-223) or inhibitors differ in their ability to respond to extracellular signals. When a growth factor or substrate capable of influencing heterodimer formation (or nuclear translocation) is identified, the experiments are repeated in other, non-myogenic cell types. The analysis of the effect of a specific growth factor in different cell types indicates whether the second secon and the second signal transduction pathway are tissue-constraints of the corresponding signal transduction pathway are tissue-constraints as a second and recording the series of the series of the expension of the contract of the contract heads were recorded and the contract of the contract o

்கு இந்த இருக்கு இருக்கு வரு படியாக studies in tissue culture cells permit the relative affinity and somether அது அருக்கு அடிய முற்ற நடிய - 12 AMACONA Manufacture of the differentiation, and subsequently in response to known signal transducers, to be evaluated to a subsequently in response to known signal transducers, to be evaluated to a subsequently in response to known signal transducers, to be evaluated to a subsequently in response to known signal transducers, to be evaluated to a subsequently in response to known signal transducers, to be evaluated to a subsequently in response to known signal transducers, to be evaluated to a subsequently in response to known signal transducers. ACCORDED To the interactions of these factors may be tested in a relevant physiological background in the second year and the competition with the prevalent endogenous components present in the cell at the time. Most analyses of the interactions of myogenic factors performed thus far have been carried out in vitro, in purified systems, or in yeast (Benezra et al., Cell, 61:1213-1230 (1990); Lassar et al., Cell, 66:305-315 (1991); Hu et al., Mol. Cell. Biol., 12:1031-1042 (1992); Hu et Chen et al., Cell, 86:731-741 (1996); and Spicer et al., Science, 272:1476-1480 (1996). The relatively low sensitivity of the biochemical methods used to directly detect interactions in mammalian cells, such as immunoprecipitation or activation of a reporter gene construct, required high levels of protein and overexpression of the construct, usually obtained by transient transfection, levels that could potentially force an interaction due to increased concentration. The methods disclosed herein permit protein-protein interactions that are functionally relevant at different points in the myogenic differentiation pathway to be studied. Clearly, the extracellular and intracellular milieu determines the stoichiometry and abundance of the these proteins at different times. As a result, competition of different proteins for the same dimerization partners, cofactors, and kinases or phosphatases in

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signal transduction pathways could have significant effects on which complexes actually form in intact cells. To assess the nature of such endogenous interactions, low expression levels are needed in order not to alter the levels inherent to the cell and characteristic of the "competitive" environment at a given time. Advantageously, high-level expression of the introduced proteins is not required in the systems described herein in order to assess the protein-protein interactions of interest. Indeed, by contrast with transient transfection assays or even most retroviral vectors with strong promoters and high translation efficiencies, the systems disclosed herein provide levels that should not perturb the natural endogenous physiological levels of the proposed test proteins in the cell:

Example 6: Analysis of inhibitory and myogenic HLH proteins in mice.

The heterodimerization of inhibitory and myogenic HLH proteins in mice may be mapped. Mtwist and I-mf have been shown to inhibit myogenesis in mammalian tissue mental state of the culture systems. In addition, they have been proposed to activia direct physical association and the contract of the culture systems. with myogenic HLH proteins (Hebrok et al., Dev. Biol., 165:537-544 (1994); Rohwedel et as a first of the second se - 15, al., Exp. Cell Res., 220:92-100 (1995); Chen et al., Cell, 86:731-741-(1996); Spicer et al., Control of the control of t Science, 272:1476-1480 (1996)). During embryogenesis, Mtwist is expressed throughout the epithelial somite and is later excluded from the myotome (Fuchtbauer, Dev. Dyn., 1997) which is the research expression has not been analyzed at early stages of somitogenesis, at 11.5 days post-coitum I-mf is highly expressed in the sclerotome but is excluded from the myotome (Chen et al., see Cell, 86:731-741 (1996)). Thus, based on their expression domains in the embryo, these factors are thought to be critical for spatial and temporal restriction of the myogenic program in early development.

> Further support for this hypothesis derives from analyses of myf5/lacZ embryos in which the myf5 coding region has been targeted and replaced by lacZ. Using β-gal as a marker of the myf5 expression pattern, cells expressing myf5 are detected in the presomitic mesoderm, where Mtwist is also expressed (Fuchtbauer, Dev. Dyn., 204:316-322 (1995); and Stoetzel et al., Mech. Dev. 51:251-263 (1995)), long before the onset of myogenesis (Cossu et al., Trends Genet., 12:218-223 (1996)). Later in development, myf5 and myoD are co-expressed together with Mtwist in the somite before the formation of a distinct myotome. Ott, et al., Development, 111: 1097-1107 (1991); Fuchtbauer, Dev. Dyn., 204:316-322 (1995); Stoetzel et al., Mech. Dev. 51:251-263 (1995); and Cossu et al.,

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Trends Genet., 12:218-223 (1996)). These cells do not express other detectable myogenic markers (Ott, et al., 1991). Thus, the reporter systems disclosed herein may be used to determine if the myf5 and MyoD proteins in these cells are maintained in an inactive state by interaction with Mtwist and/or I-mf in heterodimers. At subsequent stages of development, Mtwist and I-mf are expressed in most of the non-myogenic mesoderm, where the expression of myogenic factors is excluded. Smith et al., J. Cell Biol., 127:95-105 (1994); Fuchtbauer, Dev. Dyn., 204:316-322 (1995); Stoetzel et al., Mech. Dev. 51:251-263 (1995); and Chen et al., Cell, 86:731-741 (1996). Possibly Mtwist and I-mf are involved in the creation of a sharp border between the myotome and the adjacent stage. 10 for tissues at this stage. and the second

The state of the s interplacementary discan provide novel insights into the complex process of patterning during somitogenesis and association. they be the case and the Such studies are not limited to mice and can easily be performed in C. elegans, a conformation of the 15 Drosophila, Xenopus, zebrafish and other experimental organisms. To date, a decimal approximation of the control of the con been available. As a result, no definitive evidence is available as to when and where where the same as a second s ുള്ളുടെയുള്ള പ്രസ്തും during embryonic development interactions of such HLH heterodimers might occur@ത്ത് ക്രൂട്ടിയത്തിലായിരുള്ള

a table of the interactions. The reporter systems disclosed herein permit detailed studies of the interactions.

Example 7: Detection of HLH heterodimers in mouse embryos

20 The β-gal complementation assay is well-suited for the detection of protein-protein $\frac{1}{2}$ interactions in vivo. Myf5- $\Delta\alpha$, MyoD- $\Delta\alpha$ and Mtwist- $\Delta\omega$ fusion proteins may be constructed. Mediation of β-gal complementation with these fusion proteins may be tested in the course of performing the experiments described above. Using well-established transgenic technology (Thomas and Capecchi, Nature, 324:34-38 (1986); and Capecchi, Science, 244: 1288-1292 (1989)), mouse lines may be generated in which one of the myf5, MyoD or Mtwist alleles has been replaced with the corresponding fusion protein. Thus myf5- $\Delta\alpha$, MyoD- $\Delta\alpha$ and Mtwist- $\Delta\omega$ fusion proteins will be expressed under the control of their endogenous promoters. The expression of the test protein can be verified in these mice. The Mtwist- $\Delta\omega$ transgenic mouse may then be crossed with the myf5- $\Delta\alpha$, and the MyoD- $\Delta \alpha$ transgenic mouse lines, and in each case the offspring may be analyzed in order to identify those carrying both of the fusion proteins. β-gal activity should only develop in those cells of the embryo in which Mtwist- $\Delta\omega$ physically associates with the myf5- $\Delta\alpha$ or

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the MyoD- $\Delta\alpha$ fusion proteins. This analysis allows mapping when and where during embryonic development Mtwist is actually interacting with myf5 and MyoD to repress the myogenic phenotype.

Example 8: Targeting strategy and engineering of necessary constructs

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The myf5- $\Delta\alpha$ fusion protein coding sequence may be inserted into the myf5 locus so that it will be expressed under the control of the endogenous myf5 regulatory elements. A similar insertion of wild type β-gal in the myf5 locus resulting in a fusion with the ATG of myf5 has been shown to reproduce faithfully the expression pattern of the endogenous gene. The targeting construct is based on the published myf5/lacZ targeting construct (Tajbakhsh and Buckingham, Proc. Natl. Acad. Sci. USA, 91:747-751 (1994); Tajbakhsh 2 10 14 with the following differences: (1) The fusion protein contains the complete myf5 coding - Approximate the second section of the Δα β-gal. (2) The fusion protein coding sequence is followed by a configuration of the Δα β-gal. (2) The fusion protein coding sequence is followed by a configuration of the Δα β-gal. down to be a subsequence on your resistance gene flanked by FRT sites (FLP recombinase targets) of This allows the second flower and and the start of 15 to the G418 selection of ES cells that have taken up and integrated the targeting construct. (3) A target of the start of the st mouse genomic DNA: During homologous recombination, strand exchange will occur and a strand exchange will occur within the homology region and as a result the diphtheria toxin expression cassette will be asset as the first of the firs excluded following integration (Capecchi, Science, 244: 1288-1292 (1989)). Clones GAZ. resulting from random integration rather than homologous recombination retain diphtheria toxin expression and will be selected against during culture, because they will die. The surviving clones are characterized by PCR, and the appropriate integration of the construct in the myf5 genomic locus is confirmed by Southern blot.

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Subsequently, the neomycin selection cassette is removed using a modified version of a previously described technique (Fiering et al., Genes Dev., 9:2203-2213 (1995)). Briefly, a plasmid expressing a bicistronic message containing FLP recombinase, an Internal Ribosomal Entry Site (IRES) and GFP is transiently transfected into the ES cell clones. GFP positive cells are clonally sorted using the fluorescence activated cell sorter (FACS). In these cells, FLP deletes the sequences between the two FRT sites, and only the B-gal coding sequence remains in the ES cell genome. Aliquots of the sorted clones are tested for sensitivity to G418, and in the sensitive clones the accurate deletion of the neomycin cassette is confirmed by PCR and Southern blotting. This approach, which

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eliminates the selectable marker, avoids interference between the exogenous promoter driving the selectable marker and the endogenous regulatory sequences as described (Olson et al., Cell, 85:1-4 (1996)).

Targeting constructs for MyoD and Mtwist have also been described (Rudnicki et al., Cell, 71:383-390 (1992); Chen and Behringer, Genes Devel., 9:686-699 (1995)) and the relevant constructs may be produced for each. Based on these available reagents, and following the scheme proposed above for the myf5-Δα strategy, vectors to target (Chen and Behringer, Genes Devel., 9:686-699 (1995)) MyoD- $\Delta\alpha$ and Mtwist- $\Delta\omega$ fusions into the endogenous MyoD and Mtwist loci of ES cells may be constructed. In each case, an ES cell line syngeneic to the available genomic DNA homology regions in the targeting construct are used, as strain differences are known to reduce the frequency of homologous recombination. The same FLP-mediated excision methodology used for the myf5 "knock where the companion of the medical property of the second of the medical resistance markers from the second of the targeted MyoD and Mtwist loci. This "in-out" strategy ensures that the fusion protein The work 15 works coding regions are under the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and associated (1985) and the control of the endogenous regulatory elements and the control of the endogenous regulatory elements and the control of the endogenous regulatory elements are control of the endogenous regulatory elements and the control of the endogenous regulatory elements are control of the endogenous washered to the with minimal extraneous flanking DNA sequences. The contract the contract the contract to the contract to

the provided that the Example 9: Analysis of the myf5-Δα/Mtwist-Δω and Analysis of the myf5-Δα/Mtwist-Δα/ entransfer over the content of the transfer of the second of the second of the second over the second of the sec

For each construct, multiple ES cell clones are injected into blastocysts. The 20 chimeric offspring obtained upon implantation into pseudopregnant females are tested for germline transmission, and heterozygous mice are obtained. One critical control in this experiment is to confirm that the expression pattern of the "knocked-in" fusion proteins faithfully mimics that reported for the corresponding endogenous factors. For this purpose, a system allowing rapid detection of the fusion proteins is provided. A transgenic mouse strain expressing a β -gal mutant ($\Delta\mu$) capable of strong complementation with either $\Delta \alpha$ or $\Delta \omega$ is generated. $\Delta \mu$ is expressed ubiquitously from the strong chicken β actin promoter. MyoD-Δα, myf5-Δα and Mtwist-Δω transgenic mouse lines are each crossed with the $\Delta\mu$ transgenic mice. Since co-expression of any of these fusion proteins with the strongly complementing $\Delta\mu$ mutant should result in readily detectable β -gal activity, it is thus possible to follow the expression pattern of our fusion proteins by X-gal staining of the embryos.

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The Mtwist- $\Delta\omega$ mouse line is crossed with MyoD- $\Delta\alpha$ and myf5- $\Delta\alpha$ transgenic mouse lines. As heterozygous mice are used for these crosses, on average 1/4 of the embryos will be double heterozygotes. These embryos are analyzed at different time points during development by staining whole mount preparations and histological sections with X-gal. The sections also are stained with the more sensitive Fluor-X-Gal fluorescent substrate (Mohler, W. A., & Blau, H. M., Proc. Natl. Acad. Sci., USA, 93:12423-12427 (1996)), to detect those cells in which the Mtwist-MyoD or the Mtwist-myf5 interaction is a rare event and the β -gal signal is consequently lower.

The strength of this approach is that β -gal activity should only appear in cells in which the interactions described above take place in vivo. This approach allows a thorough analysis of the interplay between inhibitors and activators of myogenesis during development. In particular, it allows analysis of the correlation between co-expression and Marghes of Mark Ship, a physical interaction of two proteins as heterodimers in an in vivo setting, the developing section of two proteins as heterodimers in an in vivo setting, the developing section of two proteins as heterodimers in an in vivo setting, the developing section of two proteins as heterodimers in an in vivo setting, the developing section of two proteins as heterodimers in an in vivo setting. This is particularly important in the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of factors which, like Mtwist, are within the case of で表現することできた。 699 (1995)):and are likely to carry:out their functions through interaction with different での変することできた 可引起的 电磁电子 化氯酚 经 determination factors. ्राक्षाची , क्षाम्बर्दान्त्रं क्षाम्बर्धान्यः । वि

Appendix to the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis of the state of β-gal complementation mutants also can be extended to an analysis. I-mf. I-mf has also been implicated as a negative regulator of myogenesis in the embryo (Chen et al., Cell, 86:731-741 (1996)). Interestingly however, I-mf and Mtwist are coexpressed throughout most of the somite. It is not clear if their presence in the same cells is merely an indication of the existence of redundant mechanisms for repressing the activity of the myogenic HLH regulators or whether the two factors have distinct functions. In the first case, the prediction would be that both I-mf and Mtwist associate with the same factors. In the second case, differences and interactions with different factors should be detectable using our experimental approach.

Example 10: Analysis of protein interactions by Fluorescence-Activated Cell Sorting (FACS)

The β-gal activity of a population of C2C12 cells that were coinfected with FRAP- $\Delta\alpha$ and FKBP12- $\Delta\omega$ (as described in Examples 1 and 2) was assayed in the presence and absence of 10 ng/ml rapamycin by FACS. FACS was carried out according to methods that are well-known in the art, e.g., Nolan et al. (1988) Proc. Natl. Acad. Sci.

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USA 85:2603-2607. Using this assay, increased β-gal activity was detected in the majority of cells after 90 minutes of rapamycin treatment (Figure 6A). A range of expression levels was observed, as evidenced by the breadth of the peak of emission in the presence and absence of the drug (compare light and dark profiles in Figure 6A). This breadth is presumably due to variable efficiency of expression of each of the retroviral vectors following integration in the target cell. This inference is supported by the finding that when the 25% of cells expressing the lowest β-gal activity in the absence of rapamycin were collected (Figure 6B) and reassayed in the presence and absence of rapamycin, the treated and untreated cell populations yield non-overlapping peaks by FACS analysis, indicating a clear distinction between the treated (light peak) and untreated (dark peak) populations (Figure 6C). Thus, non-overlapping populations of cells distinguished by the expression (or non-expression) of complementing fusion proteins can be identified and and appropriate the second of the second and the second appropriate and the second appropriate and the second and the second appropriate and the second and the second and the second appropriate and the second and the

Example 11: Monitoring of EGF Receptor Dimerization in Living Cells

A previously unrecognized mode of regulation of the epidermal growth factor (EGF) receptor signaling pathway that acts through receptor dimerization was revealed using the methods of the invention for monitoring protein-protein interactions at the membrane of live cells. Chimeric proteins containing the extracellular and transmembrane domains of the EGF receptor, fused to weakly complementing β-galactosidase (β-gal) deletion mutants, were expressed in myoblasts. Treatment of the cells with EGF resulted ... in chimeric receptor dimerization as assessed by a rapid increase in β-gal enzymatic activity. Further treatment with EGF did not restimulate dimerization unless an inhibitor of EGF receptor tyrosine kinase was added. These results reveal a feedback mechanism in which tyrosine kinase activity of the dimeric receptor inhibits further dimerization of the receptor.

Methods

Construction of chimeric receptors. The weakly complementing $\Delta \alpha$ and $\Delta\omega$ deletion mutants of β -gal were each linked to a polypeptide sequence containing the extracellular and transmembrane domains of the human EGF receptor to form chimeric receptor molecules. The chimeric receptors lacked the cytoplasmic domain, and attendant tyrosine kinase activity, of the native receptor. The procedure was as follows. The sequence coding for the extracellular and transmembrane domains of the human EGF

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receptor (amino acids 1-655) was amplified by polymerase chain reaction (PCR) using primers that incorporated an NcoI site at the 5' end and an XhoI site at the 3' end of the PCR product. Although this fragment retains threonine 654, which is a site of protein kinase C (PKC) phosphorylation, arginines 656 and 657 are removed, destroying the consensus PKC recognition sequence. The amino acid sequence beginning with threonine 654 is thr-leu-glu-ser-met, with the met residue being the beginning of the β-gal sequence. The glu and ser codons are generated by the junction sequence and are not native to either EGF or β-gal.

DNAs encoding the chimeric receptors were inserted into a retroviral vector also encoding a selectable marker. For the construct containing the EGF receptor- $\Delta\alpha$ fusion, the selectable marker was the neo gene, encoding G418 resistance; while the EGF receptor-Δω fusion specified hygromycin resistance (Figure 1B). Accordingly, the EGF receptor PCR product was digested and cloned into the Ncol and Xhol sites of the war and t pWZL-Δα and pWZL-Δω vectors. The pWZL-Δα-neo and pWZL-Δω-hygro plasmids were constructed by cloning the lacZ Δα and Δω deletion mutants into pWZL-neo and pWZL-hygro, respectively. Mohler et al., supra; and Rossi et al. (1997) Proc. Natl. Acad. Sci. USA 94:8405-8410. Plasmids were transfected into oNX cells using Lipofectamine (Life Technologies), and virus-containing supernatant was harvested 48-72 hours later. C2F3 mouse myoblasts (Rastinejad et al. (1993) Cell 72:903-917) maintained in DME with 20% fetal bovine serum (FBS) in 10% CO₂, were infected by overnight incubation in the viral supernatant. Cells containing both constructs were selected in 1 mg/ml G418 plus 1 mg/ml hygromycin, and were maintained in 400 μg/ml of each antibiotic.

> Cells were treated with mouse salivary EGF treatment and FACS analysis gland EGF (Sigma) at 100 ng/ml and in some experiments were treated with tyrphostin AG1478 (Calbiochem) at 100 nM. Following all treatments, cells were rinsed with phosphate buffered saline (PBS), trypsinized, and resuspended in PBS + 5% FBS. Fluorescein di-β-D-galactopyranoside (FDG; Molecular Probes) was loaded into the cells by hypotonic shock as described. Fiering et al. (1991) Cytometry 12:291-301 and Nolan et al. (1988) Proc. Natl. Acad. Sci. USA 85:2603-2607. Cells were kept on ice until analysis on the cell sorter, which was conducted 1 to 2 hours after trypsinization.

The chimeric receptor was detected by immunofluorescence using a monoclonal mouse anti-human EGF receptor antibody diluted 1:100 (clone EGFR1, Dako) and either

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phycoerythrin-labeled horse anti-mouse IgG (Vector) or fluorescein-labeled goat antimouse IgG (Cappel) diluted 1:100. Cells were trypsinized and stained in PBS + 5% FBS. For each sample, FACS analysis data was collected for 5000 cells. Cells were cloned on a Becton-Dickinson FACS Vantage and analyzed on a Becton-Dickinson FACScan at the Stanford Shared FACS Facility. Data analysis was facilitated by FlowJo software (Tree Star. Inc.). Data shown here as FACS profiles were adjusted for autofluorescence using autofluorescence compensation. Alberti et al. (1987) Cytometry 8:114-119. Mean fluorescence data were adjusted for autofluorescence and for endogenous mammalian βgal activity by subtracting the mean fluorescence of untransduced cells loaded with EDG are and the control of th ~ 10 substrate.

Results

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Receptor dimerization assay. The two chimeric DNAs were each cloned 28/39 Speeding a week steer into retroviral speedors encoding selectable markers (Fig. 7B) and transduced into the C2F3 and the speeding selectable markers (Fig. 7B) and transduced into the C2F3 and the speeding selectable markers (Fig. 7B). சூர்களையில் சிக்கால்கள் myoblast cell line. After selection with G418 and hygromycin, β-gal-énzyment வகை வகிக்காகிய கிறைகள் activity was monitored using the fluorescence activated cell sorter (FACS) to measure the grant of the grant of the fluorescence activated cell sorter. at the substrate. In the absence of EGF, the population of a page of the population transduced cells consisted of a mixture of cells with low and high levels of βgal activity reserved and the EGF receptor is a result of the gray curve), which was not unexpected given that the EGF receptor is a result of the gray curve). capable of dimerizing in the absence of EGF. Gadella et al. (1995) J. Cell Biol. 129:1543-1558. Following stimulation of the population of cells with EGF many of the cells exhibited increased β-gal activity (Fig. 7C, dark gray curve). FACS analysis with an antibody specific to the human EGF receptor showed that the cells expressed a broad range of levels of the chimeric receptor (Fig. 7D, medium gray curve). Clones from this population were isolated and screened for low background levels of β--gal activity in the absence of EGF, and increased levels of β -gal activity in the presence of EGF. One such clone had a low level of chimeric receptor expression relative to the population (Fig. 7D, dark gray curve) and exhibited a several-fold increase in β-gal activity in the presence of EGF (Fig. 7E), indicating dimerization of the chimeric receptor. Dimerization was also observed following stimulation with other EGF-like growth factors that bind and activate the EGF receptor, such as TGF-\alpha, heparin-binding EGF-like growth factor, and betacellulin: but not with EGF-like factors, such as heregulin α, that act through related receptors other than the EGF receptor. Beerli et al. (1996) J. Biol. Chem. 271:6071-6076.

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Dimerization, expressed as the mean fluorescence or β-gal activity of the cells, could be detected with EGF treatments as short as one minute, and dimerization increased rapidly with longer exposure to EGF (Fig. 7F).

In order to follow the fate of Time-course of EGF Receptor dimerization. receptor dimers over time, cells from the same clone described above were cultured in media containing EGF for 0 to 24 hours and then analyzed by FACS. Dimerization peaked after 2 to 4 hours in EGF, and then decreased (Fig. 8). The fold increase in dimerization and the rate of the ensuing decline in dimerization differed among experiments, but the overall pattern was consistent, and was also observed with the original population of uncloned cells. By contrast, measurement of the levels of the chimeric receptor on the cell surface by immunofluorescence using the FACS showed that the amount of chimeric receptor on the cell surface remained essentially constant over the period that dimerization grand the second ANNULL FAREST CONTROL WAS due to either the depletion of EGE from the media, or to an inhibition of receptor face of the Control Research permitted in 15/particles dimerization. The electron of the appropriate and the state of the electron of the e

The 19th the least the second of the second dimerization, the response to a second EGF treatment was minimal, suggesting that the control of ்து கூறு நடித்து cells were resistant to further EGF-mediated dimerization despite the continued presence வருக்கு கண்களின்ற of the chimeric receptor on the cell surface. By contrast, if, following EGF treatment, cells 2 were incubated in media lacking EGF for several hours, dimerization could be restimulated with a second treatment of EGF. This indicated that the continued presence of EGF in the media was the basis for the continued inhibition of dimerization of the receptor. A. possible explanation for these results is that signaling through the endogenous wild-type EGF receptors in the cells inhibits dimerization of the chimeric receptor. A test of this hypothesis was possible, using AG1478, a highly specific inhibitor of the EGF receptor tyrosine kinase. Levitzki et al. (1995) Science 267:1782-1788.

> Accordingly, cells expressing chimeric receptor were treated with EGF overnight, and then retreated with EGF or tyrphostin. As shown in Figure 9A (left panel), sample I received a single overnight treatment with 100 ng/ml EGF. Samples II and III also were treated with EGF overnight, and then retreated with 100 ng/ml EGF for 2 hours (sample II), or 100 nM tyrphostin AG1478 for 2 hours (sample III). Sample IV received a single 2 hour treatment with 100 ng/ml EGF, and sample V received no treatment. The results

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(Figure 9A, right panel) show that treatment of the cells with tyrphostin led to an increase in dimerization, yielding dimerization levels that were comparable to the peak levels observed after a single two hour treatment with EGF, indicating that EGF receptor tyrosine kinase activity is involved in inhibiting receptor dimerization. Tyrphostin treatment also caused an increase in the amount of β-gal activity observed when previously unstimulated cells were treated with EGF. Cells were treated with EGF and tyrphostin, or EGF alone, over periods ranging from 0-24 hours. Cells that received both tyrphostin and EGF showed greater β-gal activity than cells that received EGF alone, for treatment times of up to 6 hours (Fig. 9B). By 8 hours of treatment, there was no difference in EGF receptor dimerization between EGF-treated cells and EGF+tyrphostin-treated cells. Repeated administration of tyrphostin every four hours did not further prolong the increased β-gal activity.

These results show that inhibition of receptor tyrosine kinase can relieve a feedback and the second second

These results also demonstrate that, using the methods and compositions of the invention, it is possible to monitor EGF receptor dimerization in live cells. They show, in addition, that receptor kinase activity is involved in regulating dimerization, the first step after ligand binding in EGF signal transduction. Dimerization is measurable following treatment of cells with EGF after as little as one minute, which indicates that the β-gal complementation is able to monitor the rapid production of newly formed protein dimers. Previous data on EGF binding, receptor internalization, and substrate phosphorylation also indicate that the receptor responds to ligand within minutes. Felder et al. (1992) J. Cell. Biol. 117:203-212; and Kiyokawa et al. (1997) J. Biol. Chem. 272:18656-18665.

Although receptor dimerization declines after a few hours, the chimeric receptor remains on the cell surface and is refractory to further dimerization in response to EGF. Inhibition of the endogenous receptor tyrosine kinase, however, permits further dimerization.

Inhibition of receptor dimerization begins immediately following receptor activation, as

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shown by the observation that including tyrphostin with the initial EGF treatment increases dimerization over the levels observed with EGF alone.

The kinetics of complementation reflect the kinetics of association of the binding The decline in EGF receptor dimerization is in contrast to partners observations using β-gal complementation to monitor the interaction of FRAP and FKBP12. See Examples 1, 2 and 10, supra; see also Rossi et al. (1997), supra. Using β-gal complementation to detect the rapamycin-mediated interaction between FRAP and FKBP12, the slowest increase in β-gal activity was seen at the earliest time points following the addition of rapamycin, but β -gal activity continued to increase for at least 20 hours. This could be due to stabilization of the chimeric protein interactions by formation of the active β-gal complex. With EGF receptor dimerization, however, the most rapid increase in B-gal activity was seen at the earliest time points after the addition of EGF to the media; whereas, after 2 to 4 hours, the β-gal activity declined. The difference between these results indicates that the dimerization kinetics observed with β-gal complementation where the second secon are not simply a reflection of β-gal complementation kinetics or stabilization, but reflect, at least to some degree, the kinetics of interaction of the non-B-gal portions of the chimeric proteins. The results also show that β-gal complementation can monitor the regulation of dimerization by other proteins.

> Comparison to previous methods Receptor dimerization has typically been studied by in vitro methods such as chemical cross-linking and immunopurification, followed by gel electrophoresis. Yarden et al. (1987) Biochemistry 26:1443-1451. Recently, EGF receptor dimerization has also been analyzed by fluorescence resonance energy transfer (FRET). Gadella et al. (1995) supra. Fluorescein and rhodamine labeled EGF was added to cells, and dimerization of the receptor was measured microscopically. Low temperature incubations and fixation of the cells was required to prevent internalization of the receptor before analysis, a problem that was avoided in the present experiments by using a non-internalizing mutant receptor. FRET can also be used to study interactions of fluorescently-labeled molecules within the cell or cell membrane; however, labeling and introduction of these molecules at sufficiently high concentration can be cumbersome. It has recently been shown that green fluorescent protein can be modified and used for FRET analysis on genetically expressed proteins. Miyawaki et al. (1997)

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Nature 388:882-887. The GFP signal, however, cannot be enzymatically amplified as is the case with β -gal.

Thus, β-gal complementation provides a rapid method for monitoring receptor dimerization in live cells. This method can be used for high throughput screening for pharmacological agents that can bind to a number of receptors and act as either agonists or antagonists. Binding data alone cannot indicate whether or not an agent can elicit a response; identifying a response, by analysis of downstream effects such as phosphorylation, involves destruction of the cells followed by in vitro analysis. β-gal complementation will also enable a screen for novel dimerization partners in a mammalian "two-hybrid" assay that, in the case of membrane receptors, can offer new insight into the regulation of signal transduction pathways.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be apparent to those skilled in the art that certain changes and modifications may be practiced. Therefore the foregoing descriptions and examples should not be construed as limiting the scope of the invention.

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CLAIMS

What is claimed is:

- 1. A reporter system component comprising: a first low-affinity reporter subunit, coupled to a first putative binding moiety; wherein the first low-affinity reporter subunit is capable of association with at least a second low-affinity reporter subunit to generate a detectable signal, said association being mediated by the first putative binding moiety.
- . · · · 2. The reporter system component of claim 1 wherein the first putative binding moiety is a protein. And the second second second second

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- ஆக்க கூற என்ற நடித்த கூறியாக The reporter system component of claim-2 wherein the protein is selected மண்ணுக்க நடித்த statistics of a signal transduction cascade, cell-surface 15 hours, receptors, proteins regulating apoptosis, proteins that regulate progression of the cell-cycle, which was transfer the large proteins involved in the development of tumors, transcriptional-regulatory proteins, we also appear to the second proteins in the development of tumors, transcriptional-regulatory proteins, we also appear to the second proteins and the development of tumors, transcriptional-regulatory proteins, we also appear to the second proteins and the development of tumors, transcriptional-regulatory proteins are the second proteins and the development of tumors, transcriptional-regulatory proteins are the second proteins and the second proteins are the second proteins and the second proteins are the translational regulatory proteins, proteins that affect cell interactions, cell adhesion molecules, proteins which are members of ligand-receptor pairs, proteins that participate in the control of the the folding of other proteins, and proteins involved in targeting to intracellular compartments.
 - A reporter system comprising the reporter system component of claim 1 and further comprising at least a second low-affinity reporter subunit coupled to a second putative binding partner of the first putative binding moiety.
 - 5. The reporter system of claim 4 wherein the binding affinity of the putative binding moieties for each other is greater than the binding affinity of the first and second reporter subunits for each other.
 - 30 6. The reporter system of claim 5 wherein the production of the signal is dependent upon the binding of the putative binding moieties.

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- 7. The reporter system of claim 4 wherein the first and second putative binding moieties are proteins.
- 8. The reporter system of claim 7 wherein the protein is selected from the group consisting of members of a signal transduction cascade, cell surface receptors, proteins regulating apoptosis, proteins that regulate progression of the cell-cycle, proteins involved in the development of tumors, transcriptional-regulatory proteins, translational regulatory proteins, proteins that affect cell interactions, cell adhesion molecules, proteins which are members of ligand-receptor pairs, proteins that participate in the folding of other proteins, and proteins involved in targeting to intracellular compartments.
- 9: The reporter system of claim 7, wherein the first and second reporter subunities each comprise a protein, and wherein said proteins are capable of associating to the subunities each comprise a protein, and wherein said proteins are capable of associating to the subunities each comprise a protein, and wherein said proteins are capable of associating to the subunities each comprise a protein, and wherein said proteins are capable of associating to the subunities each comprise a protein, and wherein said proteins are capable of associating to the subunities each comprise a protein, and wherein said proteins are capable of associating to the subunities each comprise a protein and wherein said proteins are capable of associating to the subunities each comprise a protein and wherein said proteins are capable of associating to the subunities each comprise a protein and wherein said proteins are capable of associating to the subunities each comprise a protein and the subunities each comprise each compri

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- 11. The reporter system of claim 7 wherein the first and second subunits are low affinity binding mutant subunits of a hydrolytic enzyme.
 - 12. The reporter system of claim 11 wherein the first and second subunits are low affinity binding mutant subunits of β -galactosidase.
 - 13. The reporter system component of claim 1 wherein said component comprises a fusion protein including said low affinity reporter subunit and said first putative binding moiety.
- 30 14. The reporter system component of claim 13 wherein the low affinity reporter subunit comprises a low affinity binding mutant subunit of β-galactosidase.

- 15. A nucleic acid encoding the fusion protein of claim 13.
- 16. The nucleic acid of claim 15 further comprising regulatory sequences effecting expression of the putative binding protein.

- 17. A viral vector comprising the nucleic acid of claim 15.
- A cell transformed with the nucleic acid of claim 15. 18.

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- "A cell transformed with a nucleic acid encoding the reporter system component of claim 13 and a nucleic acid encoding at least a second of said reporter. system components. 我好好看到我们也没有一个,我们还是我们的一个大人,你就就是我们的一个我们的,我们就是这个人,一个我们的一个,我们的一个我们的,我们就是我们的一个,我们的一个人
- 20. The reporter system component of claim 14 wherein the fusion protein the succession of the reporter system component of claim 14 wherein the fusion protein the succession of the second of the se 15 further comprises an additional protein sequence between said reporter subunit and said a protein sequence between said reporter subunit and said a protein sequence. TO THE STREET OF THE PURITURE BINDING MOIETY: THE AREA STREET OF THE AREA STREET, THE STREET OF THE AREA STREET, THE AREA ST

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21. A method of determining the occurrence of binding between first and second putative binding moieties, the method comprising:

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a) providing a reporter system comprising:

a first component comprising a first low affinity reporter subunit. coupled to the first putative binding moiety, and

a second component comprising a second low affinity reporter subunit coupled to the second putative binding moiety:

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wherein the first low affinity reporter subunit is capable of association with at least the second low affinity reporter subunit to generate a detectable signal, said association being mediated by the binding of the first and second putative binding moieties:

- b) combining the first component and the second component; and
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- c) detecting the presence or absence of the signal.

- 22. The method of claim 21 wherein the binding affinity of the putative binding moieties for each other is greater than the binding affinity of the first and second reporter subunits for each other.
- 23. The method of claim 21 wherein the first and second putative binding moieties are proteins.
- The method of claim 23 wherein the protein is selected from the group 24. consisting of members of a signal transduction cascade, cell surface receptors, proteins regulating apoptosis, proteins that regulate progression of the cell-cycle, proteins involved and apoptosis. in the development of tumors, transcriptional-regulatory proteins, translational regulatory proteins, proteins that affect cell interactions, cell adhesion molecules, proteins which are members of ligand-receptor pairs, proteins that participate in the folding of other proteins made that the folding of other proteins and the folding of other proteins are the folding of other proteins and the folding of other proteins are the f The state of the same proteins involved in targeting to intracellular compartments, resistant and proteins involved in targeting to intracellular compartments, resistant and proteins involved in targeting to intracellular compartments.

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the second reporter subunity each the method of claim 21, wherein the first and second reporter subunity each the second reporter subunity eac comprise a protein, and wherein said proteins are capable of associating to catalyze a the work who we reaction to produce a detectable signal. The work was the second control of the second was a second with the second control of the second

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- The method of claim 25 wherein the associated reporter subunits catalyze a 26. reaction to produce a product which is directly detectable as the detectable signal.
- 27. The method of claim 26 wherein the first and second subunits are low affinity binding mutant subunits of β-galactosidase.

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- 28. The method of claim 21 wherein each of said first and second components comprises a fusion protein.
- The method of claim 28 wherein the low affinity reporter subunit comprises a low affinity binding mutant subunit of β-galactosidase.

- 30. The method of claim 28 wherein step (a) comprises transforming a cell with one or more nucleic acids encoding the fusion proteins.
- 31. The method of claim 30 wherein step (c) comprises detecting the signal within the cell.
 - 32. The method of claim 30 wherein the one or more nucleic acids encoding the fusion proteins further comprise sequences regulating expression of the putative binding protein.

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- 33. The method of claim 30 wherein the fusion proteins are encoded by a viral vector.
- The method of claim 28 wherein the fusion protein further comprises as the second of claim 28 wherein the fusion protein further comprises as the second of the result of the fusion protein further comprises as the second of the second of the fusion protein further comprises as the second of the fusion of the fusion protein sequence between said reporter subunit and said putative binding mojety, for the second of the fusion of the fusion protein sequence between said reporter subunit and said putative binding mojety, for the second of the fusion of the fusion protein further comprises as the second of the fusion of the fusion protein further comprises as the second of the fusion of the fusion protein further comprises as the second of the fusion of the fusion protein further comprises as the second of the fusion of th

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- 35. The method of claim 21 wherein the degree of binding is quantitated.
 - 36. The method of claim 21 wherein the method further comprises detecting the effect of a third moiety on the binding of the first and second binding moieties, the method further comprising, after step (a) and prior to step (b), combining said reporter system with said third moiety.
 - 37. The method of claim 21 wherein the method further comprises determining potential agonist or antagonist activity of said third moiety.

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- 38. The method of claim 31 wherein the intracellular localization of the signal is determined.
- 39. The method of claim 21 wherein step (b) comprises combining the first and second components in the presence of a substance to determine the effect of the substance on binding of the first and second binding moieties.

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- 40. The method of claim 39 wherein the substance is a putative binding inhibitor of binding moieties having a predetermined binding affinity, and wherein the absence of the signal in step (c) provides an indicator that the substance is a binding inhibitor.
- The method of claim 39 wherein the substance is a putative promoter of binding between binding moieties having low or substantially no binding affinity for each other, and wherein the presence of the signal in step (c) provides an indicator that the substance is a promoter of binding of the binding moieties.
- 42. The method of claim 39 wherein the first and second reporter units, and first were the second binding moieties, each are proteins; The second binding moieties are proteins; The second binding mo

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wherein step (b) comprises expressing nucleic acid sequences encoding the first and second components within a cell suspected to contain the substance which inhibits or the promotes binding of the binding moieties; and

wherein step (c) comprises detecting the presence or absence of the signal in the cell or lysate thereof, thereby to determine the presence or absence in the cell of the substance which acts as an inhibitor or promoter of binding between the binding moieties.

- 43. The method of claim 39 wherein the substance is selected from the group consisting of a protein, lipid, carbohydrate, nucleic acid and a small molecule pharmaceutical.
- 44. A method of screening for binding of a first binding moiety with members of a plurality of different second putative binding moieties, the method comprising:
 - a) providing a plurality of reporter systems each comprising:
 a first component comprising a first low affinity reporter subunit coupled to
 the first binding moiety, and

one of a plurality of second components each comprising a second low affinity reporter subunit coupled to one of said plurality of second putative binding moieties, wherein in each of said second components, said second putative binding moiety is different;

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wherein the first low affinity reporter subunit is capable of association with the second low affinity reporter subunit to generate a detectable signal upon the binding of the first binding moiety with one of said different second putative binding moieties;

c) detecting the presence or absence of the signal in each of the binding assay

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individually combining the first component with each of the plurality of second components to produce a plurality of binding assay samples, each of which includes the first component and a different one of the second components; and

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15 15 45: W. The method of claim 44 wherein the first and second components each 1, 14, 14, 14, 14, 14 高い 1992 こと 1997 で comprise a fusion protein including the binding moiety and the reporter subunit. 日の またから こうか e のをかっ

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16. The method of claim 45 wherein, in step (b), the components are expressed that the state of from a nucleic acid sequence introduced into a cellar

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- The method of claim 46, wherein the plurality of second putative binding 47. moieties are encoded by members of a cDNA library.
 - The method of claim 47, wherein the cell is a eukaryotic cell. 48.

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- The method of claim 48, wherein the cell is a mammalian cell. 49.
- The method of claim 49, wherein the cell is a human cell. 50.

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The method of claim 44, wherein, in step (c), the signal is quantitated. 51.

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- 52. The method of claim 44, wherein cells in which binding between the first binding moiety and one of the plurality of putative second binding moieties has occurred are separated from cells in which said binding has not occurred.
- The method of claim 52, wherein separation is by fluorescence-activated 53. cell sorting.
- The method of claim 44, wherein the first binding moiety is selected from 54. the group consisting of cell surface receptors, transcriptional regulatory proteins, translational regulatory proteins, replication proteins, splicing proteins, signal transduction proteins, cell-cell adhesion molecules, cell-substrate adhesion molecules, cell-cycle proteins, oncogene products, tumor suppressor proteins, membrane receptors, proteins regulating apoptosis, developmental regulatory proteins, proteins that affect cells are the second and the second are the second and the second are the seco interactions, proteins that participate in the folding of other proteins, proteins involved in which is a large of the proteins and the proteins involved in which is a large of the proteins and the proteins involved in which is a large of the proteins and the proteins involved in which is a large of the proteins and the proteins are the proteins and the proteins are the proteins and the proteins are the protein 15 targeting to intracellular compartments, viral proteins and cytoskeletal proteins.
 - The method of claim 39 wherein the substance is a peptide, drug or synthetic analog. ** -5*
 - The reporter system of claim 4 wherein the first putative binding moiety 56. and the second putative binding partner comprise the same molecule.
 - A method of determining the occurrence of association between first and 57. second moieties, the method comprising:
 - providing a reporter system comprising:

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a first component comprising a first low affinity reporter subunit, coupled to the first moiety, and

a second component comprising a second low affinity reporter subunit coupled to the second moiety;

wherein the first low affinity reporter subunit is capable of association with at least the second low affinity reporter subunit to generate a detectable signal, said association being mediated by association between the first and second moieties,

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wherein association between the first and second moieties is mediated by a third moiety;

- b) combining the first component, the second component and the third moiety; and
- c) detecting the presence or absence of the signal.
- 58. The method of claim 57 wherein the association between the first and second moieties is mediated by multiple additional moieties.
- The reporter system of claim 1, wherein the low-affinity reporter subunits comprise an enzyme and an inhibitor of the enzyme.
- The method of claim 39, wherein the substance directly or indirectly affects of the first and second binding of the first and

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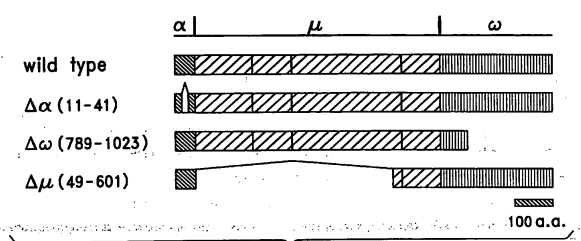
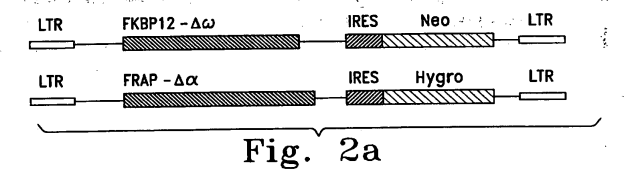
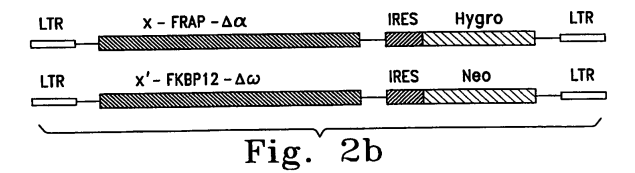


Fig. 1





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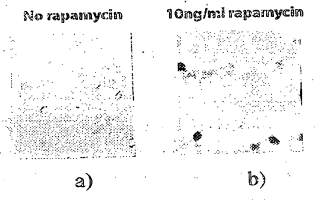
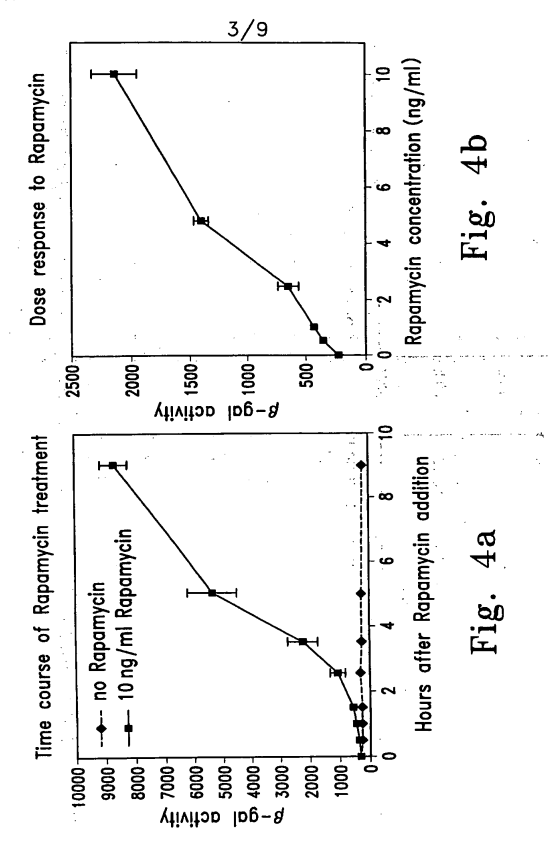


Fig. 3

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Effect of Rapamycin on lysates

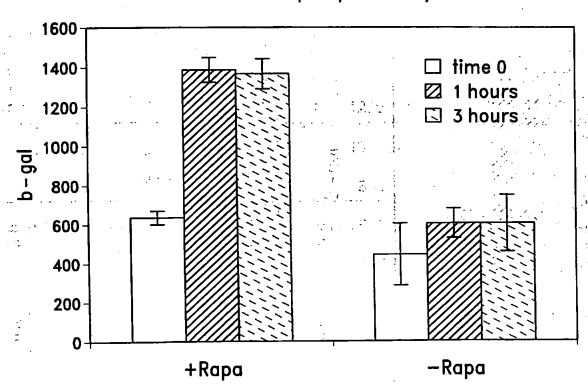
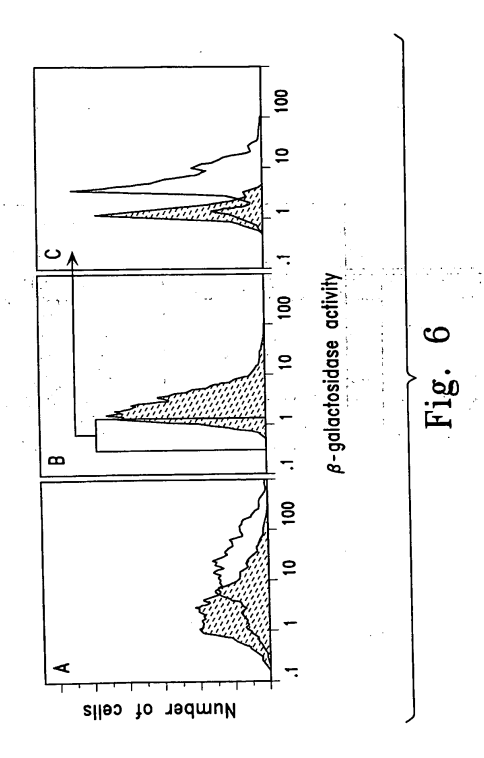
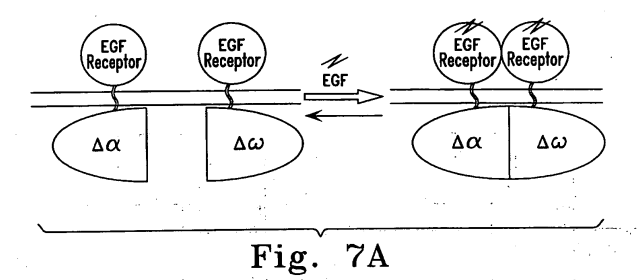


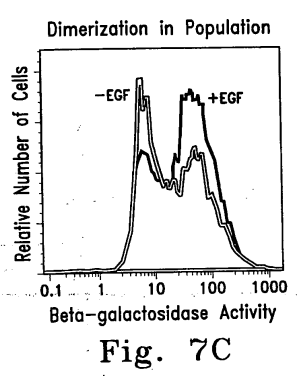
Fig. 5



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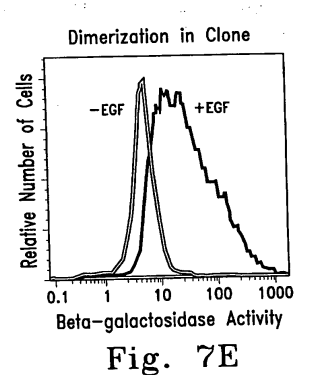
Chimeric Receptor Expression

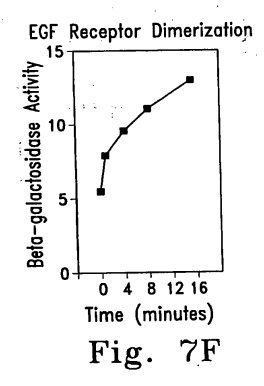
untransduced clone population

0.1 1 10 100 1000

Beta-galactosidase Activity

Fig. 7D





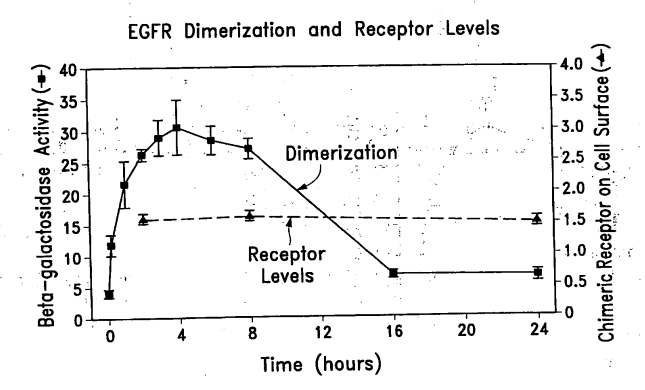
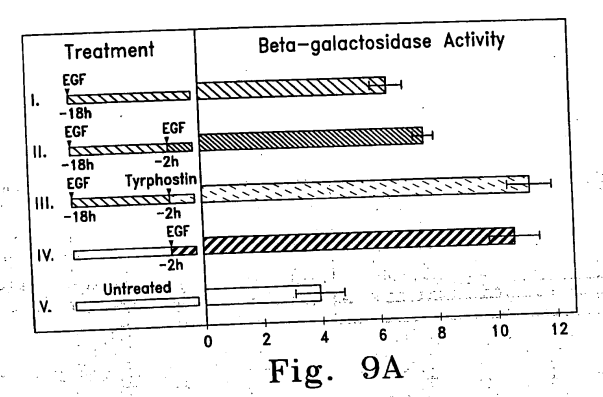
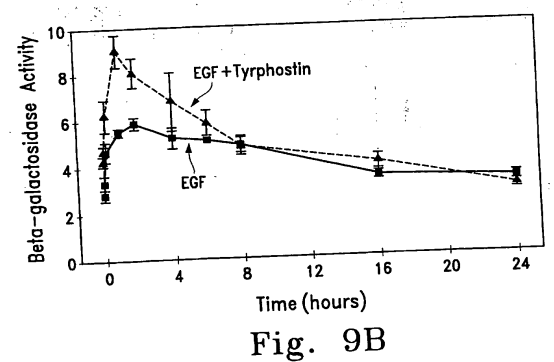


Fig. 8





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